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Psychological therapies for the management of chronic pain (excluding headache) in adults (Review)

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[Intervention Review]

Psychological therapies for the management of chronic pain (excluding headache) in adults

Amanda C de C Williams¹, Christopher Eccleston², Stephen Morley^{3a}

¹Research Department of Clinical, Educational & Health Psychology, University College London, London, UK. ²Centre for Pain Research, University of Bath, Bath, UK. ³University of Leeds, Leeds, UK

^aDeceased

Contact address: Amanda C de C Williams, Research Department of Clinical, Educational & Health Psychology, University College London, Gower Street, London, WC1E 6BT, UK. amanda.williams@ucl.ac.uk, ucjtamw@ucl.ac.uk.

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ABSTRACT

Background

Psychological treatments are designed to treat pain, distress and disability, and are in common practice. This review updates and extends the 2009 version of this systematic review.

Objectives

To evaluate the effectiveness of psychological therapies for chronic pain (excluding headache) in adults, compared with treatment as usual, waiting list control, or placebo control, for pain, disability, mood and catastrophic thinking.

Search methods

We identified randomised controlled trials (RCTs) of psychological therapy by searching CENTRAL, MEDLINE, EMBASE and Psychlit from the beginning of each abstracting service until September 2011. We identified additional studies from the reference lists of retrieved papers and from discussion with investigators.

Selection criteria

Full publications of RCTs of psychological treatments compared with an active treatment, waiting list or treatment as usual. We excluded studies if the pain was primarily headache, or was associated with a malignant disease. We also excluded studies if the number of patients in any treatment arm was less than 20.

Data collection and analysis

Forty-two studies met our criteria and 35 (4788 participants) provided data. Two authors rated all studies. We coded risk of bias as well as both the quality of the treatments and the methods using a scale designed for the purpose. We compared two main classes of treatment (cognitive behavioural therapy(CBT) and behaviour therapy) with two control conditions (treatment as usual; active control) at two assessment points (immediately following treatment and six months or more following treatment), giving eight comparisons. For each comparison, we assessed treatment effectiveness on four outcomes: pain, disability, mood and catastrophic thinking, giving a total of 32 possible analyses, of which there were data for 25.

Main results

Overall there is an absence of evidence for behaviour therapy, except a small improvement in mood immediately following treatment when compared with an active control. CBT has small positive effects on disability and catastrophising, but not on pain or mood, when compared with active controls. CBT has small to moderate effects on pain, disability, mood and catastrophising immediately post-treatment when compared with treatment as usual/waiting list, but all except a small effect on mood had disappeared at follow-up. At present there are insufficient data on the quality or content of treatment to investigate their influence on outcome. The quality of the trial design has improved over time but the quality of treatments has not.

Authors' conclusions

Benefits of CBT emerged almost entirely from comparisons with treatment as usual/waiting list, not with active controls. CBT but not behaviour therapy has weak effects in improving pain, but only immediately post-treatment and when compared with treatment as usual/waiting list. CBT but not behaviour therapy has small effects on disability associated with chronic pain, with some maintenance at six months. CBT is effective in altering mood and catastrophising outcomes, when compared with treatment as usual/waiting list, with some evidence that this is maintained at six months. Behaviour therapy has no effects on mood, but showed an effect on catastrophising immediately post-treatment. CBT is a useful approach to the management of chronic pain. There is no need for more general RCTs reporting group means: rather, different types of studies and analyses are needed to identify which components of CBT work for which type of patient on which outcome/s, and to try to understand why.

PLAIN LANGUAGE SUMMARY

Psychological therapy for adults with longstanding distressing pain and disability

Many people have pain that lasts for a long time, pain that is not relieved by drugs, surgery or physical therapy. The search for a diagnosis and for pain relief is often long, discouraging and even damaging. For some people, the pain leads to disability, depression, anxiety and social isolation. It is also associated with a tendency to experience much or all in life as ruined by pain, as a catastrophe that is impossible to control. These major life changes are not inevitable and are thought to be at least partly reversible using a treatment which aims to reduce disability and distress despite continuing pain. Treatment is based on robust psychological principles that have developed over 40 years of clinical use.

Our search found 42 trials of treatments which met our criteria, but only 35 provided data in a form that could be used. The two main types of psychological treatment are called cognitive behavioural therapy (CBT) and behaviour therapy. Both focus on helping people to change behaviour that maintains or worsens pain, disability, distress and catastrophic thinking; CBT also directly addresses the thoughts and feelings that are a problem for people with persistent pain. The effects of these two treatments on pain, disability, mood and catastrophic thinking were tested immediately after the treatment, and six months later.

Small to moderate benefits, more for disability, mood and catastrophic thinking than for pain, were found in trials which compared CBT with no treatment. Some of these were still positive six months later. Behaviour therapy showed few and only brief benefits. Psychological therapies can help people with chronic pain reduce negative mood (depression and anxiety), disability, catastrophic thinking, and in some cases, pain. Although the overall effect is positive, we do not know enough about exactly which type of treatment is best for which person.

BACKGROUND

Chronic pain is a common problem causing significant distress and disability. Behavioural and cognitive treatments designed to ameliorate pain, distress and disability were first introduced over 40 years ago and are now well established (Fordyce 1968; Keefe Rumble 2004). There are many uncontrolled trials, case studies, observations and clinical reports of treatment methods. Narrative reviews generally report positive effects of psychological treatments on a range of outcomes. In addition there has been periodic

publication of meta-analyses and systematic reviews (Flor 1992; Morley 1999) and many recent studies have focused on specific patient groups such as those with musculoskeletal pain syndromes (Dixon 2007; Guzman 2001; Hoffman 2007; Henschke 2010a), and older adults (Ersek 2008).

There is a broad family of treatments included in the general term 'psychological'. In essence, treatments have been developed that are specifically designed to alter psychological processes thought to underlie or significantly contribute to pain, distress and/or disability. The design of psychological treatments is normally informed by specific theories of the aetiology of human behaviour, or treatments have developed pragmatically through observation and study of response to intervention. In practice there is variety in the types of interventions used, and not all have been evaluated for their effectiveness. The evidence base for psychological therapies is dominated by studies of programmatic and protocolised treatments from a behavioural or cognitive behavioural tradition of clinical psychology. Psychological therapies are commonly presented as being offered after orthodox treatments have failed, when the treatment goal shifts from one of removing or alleviating pain to one of managing pain and its myriad adverse consequences on quality of life. A typical treatment protocol for cognitive behavioural therapy (CBT) will involve methods aimed directly at assessing the thoughts associated with pain, the extent of avoidance of unpleasant thoughts and of painful experiences, and the consequences of these. A common focus is on strongly held beliefs about pain and their relationship with behaviour, which typically worsens the situation in the shorter or longer term. Behavioural methods focus on the identification of behaviour that is contingent on pain, or upon events which provide pain relief or comfort, and the development of behaviour that is contingent instead on goal achievement related to the values of the individual with pain. Most therapies involve education, and many are incorporated within larger treatment programmes involving physical and occupational therapy.

In earlier reviews on this topic (Eccleston 2009a; Morley 1999), we searched for all published randomised controlled trials (RCTs) of interventions described as psychological in nature, and recovered trials principally of behaviour therapy or CBT (Morley 1999). RCTs of interventions for headache were excluded for several reasons: for consistency with the previous review (Eccleston 2009a); and because CBT for headache aims primarily at reducing frequency, duration and intensity of headache pain rather than at rehabilitating despite ongoing pain. Readers are referred to other reviews (Nestoriuc 2007; Nestoriuc 2008; Nicholson 2004), although there are no recent systematic reviews. The Eccleston 2009a review found 52 trials, of which 40 had data that could be entered into a meta-analysis. Trials of CBT provided more data than did behaviour therapy, particularly in relation to active controls. Against active control, CBT improved disability post-treatment, and pain, disability and mood at follow-up, although effect sizes were small. Surprisingly, against doing nothing (treatment as usual or waiting list control), there was only significant improvement for pain post-treatment and mood at follow-up. Again, effect sizes were small. Compared with doing nothing, behaviour therapy improved pain post-treatment, but showed no other benefits, and there were too few trials of behaviour therapy against active control for analysis. This analysis is now out of date and in need of updating (Shojania 2007). Other developments in psychological science have led to new forms of treatments being promoted, and the quality of trials and trial reporting is thought to be improving (Morley 2006). The aim of this review is to summarise the published evidence on the efficacy of psychological treatments for chronic pain in adults and, as far as possible, to investigate key variables that are thought to influence the effectiveness of many psychological interventions.

OBJECTIVES

To determine the clinical effectiveness of psychological therapy for non-malignant chronic pain (excluding headache) for adults compared with medical or physical treatments, placebo or waiting list controls.

METHODS

Criteria for considering studies for this review

Types of studies

RCTs comparing a credible psychological treatment, or a compound treatment with primary psychological content, with placebo, other active treatment, treatment as usual, or waiting list control, in chronic pain. Studies were excluded if they were concerned with headache or associated with a malignant life-threatening disease. We judged a psychological treatment credible if it was based on an extant psychological model or framework, and its delivery was from, or was supervised by, a healthcare professional qualified in psychology.

Studies were included if they:

- were available as a full publication or report of a RCT;
- had a design that placed a psychological treatment as an active treatment of primary interest;
- had a psychological treatment with definable psychotherapeutic content;
- were published (or electronically pre-published) in a peerreviewed science journal;
- were with participants reporting chronic pain (i.e. at least three months' duration); and

• had 20 or more participants in each treatment arm at the end of the treatment assessment.

This last criterion of $N \ge 20$ at post-treatment assessment is an improvement from the Eccleston 2009a review in which we used an entry of $N \ge 10$. We made this change because of the recognised risk of bias of small numbers (Ioannidis 2005; Nuesch 2009); raising the required N further would be desirable but would exclude too many studies.

Types of participants

Adults (aged 18 years or older) reporting pain of at least three months' duration in any body site, not associated with a malignant disease process. Patients with only headache or migraine were excluded because the psychological treatments for headache and migraine are sufficiently different, and have a separate history (see Nestoriuc 2007; Nestoriuc 2008; Nicholson 2004), although an up to date systematic review is lacking.

Types of interventions

Studies were included if at least one trial arm consisted of a psychology intervention, with at least one comparator arm of a placebo condition, other active treatment, treatment as usual or waiting list control.

Types of outcome measures

- We collected data on descriptive characteristics of participants and characteristics of the treatments, including treatment setting, mode of delivery and therapist.
- Following the Eccleston 2009a review, we collected data for this review on outcomes in the domains of pain experience, disability, negative mood and catastrophic thinking; we recorded and described all outcomes.

Search methods for identification of studies

Electronic searches

We identified RCTs of any psychological therapy in the Cochrane Central Register of Controlled Trials (CENTRAL 2011, issue 3), MEDLINE, EMBASE and Psychlit from their inception to September 2011. We identified additional studies from the reference lists of retrieved papers and from discussion with investigators. We performed searching in two sets. We undertook the first prior to the previously published systematic review (Morley 1999). We undertook the second focusing on the 10 years since that review using the same search strategy but taking account of changes in search architecture and terminology (see Eccleston 2009a). There were two further searches to update: in December 2009 covering

the period from the beginning of abstracting services to December 2009, and in October 2011, covering the period from December 2009 to September 2011. The search sampled the same databases; an example search strategy is given in Appendix 1. We applied no language restrictions. At least two review authors reviewed all abstracts and they were included on the basis of consensus agreement and discussion with the third review author when necessary.

Data collection and analysis

Selection of studies

The trials used in the previous systematic review and meta-analysis (Eccleston 2009a) were automatically included, although some were subsequently excluded by the stricter criteria adopted here. The two searches of the literature since the end of the previous search produced a set of possible abstracts. From these, one rater selected for examination all full papers which might meet the criteria. All three authors read the papers and agreed on exclusion or inclusion: we rated the final set of papers, including those eligible from the previous systematic review, for quality and extracted data.

Data extraction and management

We used a data extraction book devised jointly by the review authors and used in the previous review (Eccleston 2009a) to extract information on the design of the study, the participants, primary diagnosis, method of treatment and outcome measurement tools used.

The primary data type was measurement using continuous scales. We estimated treatment effects using standardised mean differences by extracting means, standard deviations and sample size at post-treatment and follow-up. When data were not available from published studies or from authors, we did not infer any parameters. Dichotomous outcome data based on clinical improvement were rare and we did not extract these.

Assessment of risk of bias in included studies

We assessed risk of bias using the recommended Cochrane guidance (Higgins 2011). Of the five suggested 'Risk of bias' categories, we included random sequence generation (selection bias), allocation concealment (selection bias), blinding of outcome assessment (detection bias), incomplete outcome data (attrition bias) and selective reporting (reporting bias). We excluded the option of 'blinding participants and personnel' because neither therapists nor patients can be blinded to whether they deliver or receive treatment. As in the previous review (Eccleston 2009a), we applied a quality rating scale specifically designed for psychological interventions in pain (Yates 2005). Two of the three review authors

scored all studies and they reached a consensus after initial comparison or ratings. The quality rating scale was designed specifically for application to psychological treatment studies in pain. It provides an overall total score (0 to 35) consisting of two subscales: a treatment quality scale (0 to 9) covering stated rationale for treatment, manualisation, therapist training and patient engagement; and a design and methods scale (0 to 26) covering inclusion/exclusion criteria, attrition, sample description, minimisation of bias (randomisation method, allocation bias, blinding of assessment, equality of treatment expectations), selection of outcomes, length of follow-up, analyses and choice of control. The first four 'Risk of bias' items from the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011) are represented in the design section of the Yates 2005 scale, accounting for up to five of the nine points available.

Measures of treatment effect

We investigated two classes of psychological treatment and labelled these cognitive behavioural therapy (CBT) and behavioural therapy. CBT involves treatments that include specific direct cognitive therapeutic content. Behavioural therapy includes treatments that are purely behavioural technologies such as biofeedback. Two classes of comparator treatments are investigated and labelled active control and treatment as usual. The active comparator involves a treatment designed to change pain behaviour such as physical therapy, education or medical regime. Patients randomised to the active control within each trial all receive the same treatment. For patients assigned to a waiting list, trials vary in whether they provide further care, and patients vary in whether they seek further care. For patients assigned to treatment as usual, this treatment can consist of anything from regular consultations to access to care. Thus patients in these conditions receive variable and usually unrecorded treatment.

Where a trial had more than two arms, we selected those which best matched our requirements for CBT or behavioural therapy, and where there was a choice, the most intensive version of either: for example, if a trial had an enriched CBT (that is, CBT with additional non-core components such as vocational guidance), a minimum CBT and a waiting list condition, we compared the enriched CBT with the waiting list. If both of the treatment conditions were eligible and fell into different analyses, each was compared with the control condition: for example, a trial comparing CBT with behavioural therapy with waiting list control was used both as CBT versus waiting list control, and behavioural therapy versus waiting list control.

We also selected two assessment time points: post-treatment and follow-up. Post-treatment is the assessment point immediately following treatment, and follow-up is the assessment point at least six months after the end of treatment, but not more than 12 months, and the longer of the two if there were two follow-up assessments within this timeframe. Therefore eight separate comparisons were

designed comprising two classes of psychological treatment under investigation (CBT, behavioural therapy), two forms of comparator (active control, treatment as usual), and two assessment time points (post-treatment and follow-up). They are labelled:

- 1. cognitive behavioural versus active control post-treatment;
- 2. cognitive behavioural versus active control follow-up;
- 3. cognitive behavioural versus treatment as usual post-
- 4. cognitive behavioural versus treatment as usual follow-up;
- 5. behavioural versus active control post-treatment;
- 6. behavioural versus active control follow-up;
- 7. behavioural versus treatment as usual post-treatment;
- 8. behavioural versus treatment as usual follow-up.

Multiple measurement tools are typically used in each trial. For each comparison we identified four outcomes and labelled them 'pain', 'disability', 'mood' and 'catastrophic thinking'. Although standard trial reporting guidance promotes the definition of primary outcomes (Boutron 2008), most trials do not state a single or preferred a priori primary outcome, so a judgement must be made. From each trial we selected the measure considered most appropriate for each of the three outcomes. When there was more than one measure for an outcome we gave preference to the measure that has documented frequent usage in the field as opposed to a novel measure. Also, when there was a choice between singleitem and multi-item self report tools, we chose longer tools on the basis of inferred increased reliability. Not all trials reported data on all three outcomes of pain, disability and mood, and not all trials reported follow-up data.

Assessment of heterogeneity

We assessed heterogeneity according to the standard method using the Chi² test and the I² statistic, calculated for each comparison on each outcome. I² values above 50% indicate high heterogeneity, between 25% and 50% medium heterogeneity, and below 25% low heterogeneity.

RESULTS

Description of studies

Results of the search

The results of the two update searches, in December 2009 and in October 2011, are described separately below.

From the 52 trials which met inclusion criteria in the original review (Eccleston 2009a), 10 trials were dropped. Eight had insufficient psychotherapeutic content, decided following further discussion of what constituted psychotherapeutic content: Astin 2003;

Becker 2000; Carson 2005; Dworkin 1994; Dworkin 2002b; Fairbank 2005; Freeman 2002; Strong 1998; one (Turner-Stokes 2003) was a trial to test equivalence of two psychological treatments and therefore on reconsideration did not meet our criteria; one (Buhrman 2004) was the only internet trial, and in the intervening period a separate review of internet interventions had been published (Macea 2010) which made it preferable to exclude internet trials from this systematic review. We included four papers which had been excluded previously: Alaranta 1994 and Spence 1995 (which with redefinition of psychological content met the criteria); Keefe 2004 (wrongly excluded for no nonpsychological comparator); and Peters 1990 (which has one outcome, N < 10). We made renewed efforts to obtain analysable data from six of the 52 studies which had not provided analysable data for Eccleston 2009a. These were Buckelew 1998; Geraets 2005; Marhold 2001; Parker 1988; Smeets 2006 and Strauss 1986. We obtained analysable data from Geraets 2005; Marhold 2001 and Smeets 2006.

The search in December 2009 produced 21 studies. Twelve studies were eligible: Babu 2007; Bliokas 2007; De Souza 2008; Ersek 2008; Falcao 2008; Leeuw 2008; Lindell 2008; Linton 2008; Morone 2008; Wicksell 2008; Woods 2008; Zautra 2008. We also found one long-term follow-up of an existing study follow-up: Smeets 2009. Eight new trials were excluded: inadequate psychotherapeutic content (Kroenke 2009; Machado 2007); internet trial (Lorig 2008); hypnosis trial (Abrahamsen 2008; Castel 2009); unclear randomisation (Ferrari 2006); inadequate N (Menzel 2006); and one which was only a trial plan (Garcia-Campayo 2009). We decided to exclude hypnosis since it fell short of classification as cognitive or behavioural treatment, and requires a systematic review devoted to it. We sought data accessible for analysis from authors and obtained data from Babu 2007; Bliokas 2007 and Zautra 2008.

The search in October 2011 produced 27 studies, of which we eventually included seven: Ehrenborg 2010; Liedl 2011; Litt 2009; Schmidt 2011; Thorsell 2011; Van Koulil 2010; and Wetherell 2011, and a further eligible study Glombiewski 2010b was not found by the electronic search but through an ineligible paper, Glombiewski 2010a, which was produced by that search). Of the 20 excluded studies, 14 had insufficient psychotherapeutic content (Carson 2010; de Sousa 2009; Dufour 2010; Esmer 2010; George 2008; Kapitza 2010; Lamb 2010; Lambeek 2009; Li 2006; Morone 2009; Rendant 2011; Sahin 2011; Turner 2011; Wong 2011); three used hypnosis (Abbott 2010; Abrahamsen 2008; Jensen 2009), one was a non-inferiority trial (Jensen 2009b); one included some participants without chronic pain (Christiansen 2010); and one was not randomly allocated (Schulze 2008). We requested missing data from authors but obtained none for included studies.

This process provided a total of 65 RCTs: Alaranta 1994; Altmaier 1992; Babu 2007; Basler 1997; Bliokas 2007; Bradley 1987; Buckelew 1998; Cook 1998; De Souza 2008; Ehrenborg 2010;

Ersek 2003; Ersek 2008; Evers 2002; Falcao 2008; Flor 1993; Geraets 2005; Glombiewski 2010b; Greco 2004; Haldorsen 1998; Hammond 2001; Jensen 1997; Jensen 2001; Johansson 1998; Kaapa 2006; Keefe 1990; Keefe 1996; Keefe 2004; Kole-Snijders 1999; Kraaimaat 1995; Leeuw 2008; Liedl 2011; Lindell 2008; Linton 2008; Litt 2009; Marhold 2001; McCarberg 1999; Mishra 2000; Moore 1985; Newton-John 1995; Nicassio 1997; O'Leary 1988; Parker 2003; Peters 1990; Puder 1988; Radojevic 1992; Redondo 2004; Schmidt 2011; Smeets 2006; Spence 1989; Spence 1995; Strauss 1986; Thieme 2003; Thorsell 2011; Turner 1988; Turner 1990; Turner 1993; Turner 2006; Van Koulil 2010; Vlaeyen 1995; Vlaeyen 1996; Wetherell 2011; Wicksell 2008; Williams 1996; Woods 2008; Zautra 2008. Of these, eight did not have analysable data: Alaranta 1994; Buckelew 1998; De Souza 2008; Kole-Snijders 1999; Lindell 2008; O'Leary 1988; Parker 1988: Strauss 1986.

We then applied the new criterion requiring N ≥ 20 in each arm of a comparison and this excluded 23 trials: Babu 2007; Bradley 1987; Cook 1998; Ersek 2003; Flor 1993; Johansson 1998; Keefe 2004; Liedl 2011; Linton 2008; Marhold 2001; Moore 1985; Newton-John 1995; O'Leary 1988; Peters 1990; Radojevic 1992; Redondo 2004; Spence 1989; Spence 1995; Turner 1990; Turner 1993; Vlaeyen 1995; Wicksell 2008; Woods 2008. We therefore proceeded with 42 trials for the review; of these, seven provided no data: Alaranta 1994; Buckelew 1998; De Souza 2008; Kole-Snijders 1999; Lindell 2008; Parker 1988; Strauss 1986.

Included studies

Sixteen of the 42 studies are new since the review of 2009 (Eccleston 2009a), meaning that more trials have been published since 2000 than before it. Of the 42 included studies, 24 had two arms, 14 had three and four had four arms. As in the 2009 systematic review, we scored the quality of trial design and found a mean 15.8/26 (standard deviation (SD) 4.3, range 9 to 24/26) which increased with year of publication (Spearman's rho = 0.41, P < 0.01); this represented an improvement of about two points per decade. The total number of patients providing data immediately post-treatment was 4788 at the end of treatment (a mean of 114 per study, SD 71) from the 5424 patients starting treatment (data from 41 of the 42 trials). Mean study completion rate from entry to post-treatment assessment was 87.6% (SD 9.5%) and ranged from 65% to 100%. Overall, the mean number of patients per trial, 114 in this review, was an increase on the mean of 91 in the 2009 review (Eccleston 2009a), although unlike the 2009 review (which included studies with N between 10 and 19), sample size did not increase with publication date. Women usually outnumbered men, with the average proportion of women per trial being 71% (SD 21%, range 4% to 100%). The mean age was 48 (SD 9, range of means from 31 to 82 years), and the mean years of pain (from the 30 studies which provided data) was 8.3 (SD 4.3, range of means from 1.3 to 16.5 years).

Forty-one of the studies specified the source of participants, who were recruited mainly from a range of healthcare settings: 16 studies recruited from pain rehabilitation clinics, one of which supplemented its participants with volunteers; two further studies drew on referrals for pain management and rehabilitation, and one study drew on dental clinic patients and volunteers (46% studies altogether recruited through pain services). Nine studies recruited from rheumatology clinics, one of which supplemented its participants with volunteers (21% altogether). Seven studies recruited from the community (including one retirement home), with an additional three community recruitment studies adding volunteers (24% studies altogether recruited from community sources), and one study recruited entirely through advertisement for volunteers. Two studies took referrals from work-based healthcare services. Nine studies (21%) were solely for patients with low back pain, and a further one for low back or neck pain; two were for spinal pain, one for neck and shoulder and one for shoulder alone; eight (19%) were for mixed chronic pain patients in which back pain was usually the most common complaint. Seven studies had patient groups with rheumatoid arthritis including one with systemic lupus erythematosus; eight had fibromyalgia; three had temporomandibular joint pain; two had osteoarthritis of the knee.

We classified treatment arms on the basis of their content and of the label given by the authors as cognitive behavioural treatment or as behavioural treatment. All treatment involved a psychologist, trained, or in training and supervised, in delivery. The mean quality of treatment was 5.4/9 (SD 2.3, range 1 to 9) and was unrelated to year of publication (Spearman's rho = 0.20, non-significant). We classified control conditions as 'active control' when there was a protocolised treatment which engaged the patient, such as an exercise programme, a medical procedure, an education programme, a support group or a self instruction booklet, and as 'waiting list or treatment as usual'. We did not distinguish between waiting list and treatment as usual because for some patients treatment as usual is elective treatment which may be none at all and therefore equivalent to being on a waiting list; and some studies allow patients on waiting lists to seek other treatment elsewhere, treatment which may be equivalent to that in 'treatment as usual' conditions. We are aware that this is not an entirely satisfactory classification where treatment as usual involves some active and regular physiotherapy or pharmacotherapy, not dissimilar to those offered in active controls, and where the large majority of patients follow it routinely, but when available information did not allow us to assign this condition to an active control, we classified a condition as treatment as usual.

Excluded studies

Ninety-three studies did not meet the inclusion criteria and were excluded. Disregarding those which did not primarily concern chronic pain, or which did not appear to be randomised, which were non-inferiority trials, which had too small a number of par-

ticipants post-treatment, or which were trials of hypnosis or internet interventions, 36 initially appeared to be trials of CBT or behavioural therapy, but on reading the full paper failed our criteria for credible psychological treatment (Abbott 2010; Appelbaum 1988; Astin 2003; Becker 2000; Bendix 1997; Broderick 2004; Brox 2003; Carson 2005; Carson 2010; de Sousa 2009; Dufour 2010; Dworkin 1994; Dworkin 2002a; Dworkin 2002b; Esmer 2010; Fairbank 2005; Fors 2000; Freeman 2002; George 2008; Haugstad 2006; Kapitza 2010; Keller 2004; Kroenke 2009; Lamb 2010; Lambeek 2009; Li 2006; Machado 2007; Moffett 2005; Morone 2009; Rendant 2011; Sahin 2011; Schweikert 2006; Soderlund 2001; Strong 1998; Turner 2011; Wong 2011). While the initial inclusion of these studies from the search is in part evidence of the diversity of terminology used to describe pain and treatments, it also raises important issues about nonspecific or design features which potentially undermine the content or fail to deliver what is implied by the description of treatment, and about the inevitably blurred boundaries between psychological intervention and education, instruction or nonspecific support. This judgement was difficult to apply in some cases and led to extended discussion between the review authors to reach a decision.

Risk of bias in included studies

'Risk of bias' is shown in Figure 1 and Figure 2: we used five 'Risk of bias' categories: random sequence generation (selection bias), allocation concealment (selection bias), blinding of outcome assessment (detection bias), incomplete outcome data (attrition bias) and selective reporting (reporting bias). Fifteen studies described a convincing method of randomisation so we judged them to have a low risk of bias, and a further 11 provide an inadequate description so we judged them to be unclear. We judged 16 to have high risk of bias, mainly because the method of randomisation was not described; these were mainly earlier studies. We judged 14 studies to have adequate allocation concealment, one uncertain and 27 high risk, again mainly because there was no description of any procedure designed to do so. Only 12 studies reported attrition fully, including finding no difference between dropouts and completers, and we judged them to have low risk of bias; 19 were unclear risk, mainly because of lack of testing for differences between dropouts and completers, but in some cases because those differences were found; and we judged 11 to have high risk of bias, predominantly because they provided no details of attrition. We judged 34 studies at low risk of bias for selective reporting of outcome since they reported all outcomes, or in one case accounted for those they did not report; we judged one study uncertain because outcomes were combined in factor scores, and seven studies did not report all outcomes which they described in assessment sections of their Methods, and we judged them at high risk of bias. Finally, we judged 13 studies at low risk of bias for outcome assessment since they used blinded assessors; two were unclear; and we judged 27 at high risk of bias since they gave no details of outcome assessment procedures. It should be borne in mind, however, that almost all outcomes were assessed by self report, so that there were restricted opportunities for influencing patients' scores. Thus most judgements of high risk of bias were because of inadequate reporting: we recognise that this is a conservative position and that some studies may have exercised proper precautions in some or all of these areas.

Figure 1. 'Risk of bias' summary: review authors' judgements about each methodological quality item for each included study.



Random sequence generation (selection bias)

Allocation concealment (selection bias)
Incomplete outcome data (attrition bias)

Selective reporting (reporting bias)

Blinding of outcome assessment (detection bias)

0% 25% 50% 75% 100%

Low risk of bias

Unclear risk of bias

Figure 2. 'Risk of bias' graph: review authors' judgements about each methodological quality item presented as percentages across all included studies.

The comprehensive quality assessment scale (Yates 2005) is reported in Characteristics of included studies. For the 42 studies which met the inclusion criteria, the mean overall quality of the studies was 21.2 (SD 5.9, range 10 to 32). The mean design quality score was 15.8 of a possible 26 (SD 4.3, range 9 to 24). A Spearman's correlation to investigate the association between year of study and overall quality score showed a weak relationship (rho = 0.37, P < 0.05), and between year of study and design quality score a slightly stronger relationship (Spearman's rho=0.41, P < 0.01). Treatment quality was **not** associated with year of study: see Included studies. N at the end of treatment was associated with design quality score and with total quality score (rho = 0.41 (P < 0.01) and 0.38 respectively (P < 0.05)).

Of the 24 analyses reported (CBT or behaviour therapy versus active control or treatment as usual, post-treatment and follow-up, for 'pain', 'disability', 'mood' and 'catastrophic thinking'), 10 showed low heterogeneity ($I^2 \le 25\%$), six showed modest heterogeneity ($I^2 > 25\%$ to < 50%) and eight, almost all analyses of behaviour therapy, showed large heterogeneity ($I^2 > = 50\%$).

Effects of interventions

Cognitive behavioural versus active control posttreatment

We entered 13 studies with 1258 participants into an analysis of the effects of cognitive behavioural therapy (CBT) on pain compared to active control. The overall effect of CBT on pain was not significant (Z = 1.43, P > 0.05) (Analysis 1.1). We entered 12 studies with 1130 participants into an analysis of the effects of CBT

on disability. The overall effect was significant (Z = 2.66, P < 0.01) with a small effect size: standardised mean difference (SMD) -0.19 (95% confidence interval (CI) -0.33 to -0.05) (Analysis 1.2); the I² value was 25%. We entered 13 studies with 1256 participants into an analysis of the effects of CBT on mood; the overall effect was not significant (Z = 0.72, P > 0.05) (Analysis 1.3). We entered six studies with 735 participants into an analysis of the effects of CBT on catastrophising; the overall effect of CBT was just significant: Z = 1.92, P = 0.05 (Analysis 1.4). The effect size was SMD -0.18 (95% CI -0.36 to 0.00) and the I² value was 31%.

Cognitive behavioural versus active control at followup

We entered 11 studies with 1261 participants into an analysis of the effects of CBT on pain at follow-up. The overall effect of CBT was not significant (Z=1.12, P>0.05) (Analysis 2.1). We entered 12 studies with 1295 participants into an analysis of the effects of CBT on disability at follow-up. The overall effect of CBT at follow-up was significant (Z=2.28, P<0.05) with a small effect size of SMD -0.15 (95% CI -0.28 to -0.02) (Analysis 2.2); the I^2 value was 23%. We entered 11 studies with 1261 participants into an analysis of the effects of CBT on mood at follow-up. The overall effect of CBT was not significant (Z=1.15, P>0.05) (Analysis 2.3). We entered two studies with 282 participants into an analysis of the effects of CBT on catastrophising. The overall effect of CBT was not significant: Z=0.49, Z=0.050 (Analysis 2.4).

Cognitive behavioural versus treatment as usual posttreatment

We entered 16 studies with 1148 participants into an analysis of the effects of CBT on pain. The overall effect of CBT was significant (Z = 2.59, P < 0.05) with an effect size of SMD -0.21 (95% CI -0.37 to -0.05) (Analysis 3.1); the I² value was 45%. We entered 15 studies with 1105 participants into an analysis of the effects of CBT on disability. The overall effect was significant (Z = 2.35, P < 0.05) (Analysis 3.2). The effect size was SMD -0.26 (95% CI -0.47 to -0.04); the I² value was 67%. We entered 12 studies with 899 participants into an analysis of the effects of CBT on mood. The overall effect of CBT was significant (Z = 3.84, P < 0.01) (Analysis 3.3). The effect size was SMD -0.38 (95% CI -0.57 to -0.18); the I² value was 49%. We entered five studies with 308 participants into an analysis of the effects of CBT on catastrophising. The overall effect of CBT was significant: Z = 4.58, P < 0.01 (Analysis 3.4). The effect size was SMD -0.53 (95% CI -0.76 to -0.31) and the I2 value was 0%.

Cognitive behavioural versus treatment as usual at follow-up

We entered seven studies with 635 participants into an analysis of the effects of CBT on pain at follow-up. The overall effect of CBT was not significant ($Z=0.99,\,P>0.05$) (Analysis 4.1). We entered six studies with 450 patients into an analysis of the effects of CBT on disability at follow-up. The overall effect of CBT was not significant ($Z=0.66,\,P>0.05$) (Analysis 4.2). We entered seven studies with 637 patients into an analysis of the effects of CBT on mood at follow-up. The overall effect of CBT was just significant ($Z=1.99,\,P=0.05$) with a small effect size of SMD - 0.26 (95% CI -0.51 to 0.00) (Analysis 4.3); the I² value was 58%. There was only one study of 59 participants in the analysis of the effects of CBT on catastrophising. The overall effect of CBT was not significant: $Z=0.84,\,P>0.05$ (Analysis 4.4).

Behavioural versus active control post-treatment

There are insufficient studies in this comparison for meta-analysis. One study of 39 participants was analysed for the effects of behaviour therapy on pain. The overall effect of behaviour therapy was not significant (Z=0.77, P>0.05) (Analysis 5.1). We entered two studies of 110 participants into an analysis of the effects of behaviour therapy on disability. The overall effect was not significant (Z=1.46, P>0.05) (Analysis 5.2). There was only one study, with 71 participants, in the analysis of the effects of behaviour therapy on mood (Analysis 5.3), with an effect of behaviour therapy that was just significant (Z=1.94, P=0.05). The effect size was SMD -0.47 (95% CI -0.94 to 0.00). We entered two studies with 146 participants into the analysis of the effects of behaviour therapy on catastrophising. The overall effect was not significant: Z=1.67, P>0.05 (Analysis 5.4).

Behavioural versus active control at follow-up

There are insufficient studies in this comparison for meta-analysis. There was only one study with 73 participants in the analysis of the effects of behaviour therapy on pain at follow-up. The overall effect of behaviour therapy was not significant (Z = 0.13, P > 0.05) (Analysis 6.1). We entered two studies with 144 participants into an analysis of the effects of behaviour therapy on disability at follow-up. The overall effect of behaviour therapy was not significant (Z = 1.01, P > 0.05) (Analysis 6.2). We entered only one study with 71 participants into the analysis of the effects of behaviour therapy was not significant (Z = 1.55, Z = 0.05) (Analysis 6.3). We entered one study with 73 participants into the analysis of the effects of behaviour therapy on catastrophising. The overall effect was not significant: Z = 0.25, Z = 0.050 (Analysis 6.4).

Behavioural versus treatment as usual posttreatment

We entered five studies of 484 participants into an analysis of the effects ofbehaviour therapy on pain. The overall effect of behaviour therapy was not significant ($Z=1.05,\ P>0.05$) (Analysis 7.1). We entered five studies of 504 participants into an analysis of the effects of behaviour therapy on disability. The overall effect was not significant ($Z=1.40,\ P>0.05$) (Analysis 7.2). We entered three studies of 278 participants into an analysis of the effects of behaviour therapy on mood. The overall effect of behaviour therapy was not significant ($Z=1.18,\ P>0.05$) (Analysis 7.3). We entered three studies with 269 participants into the analysis of the effects of behaviour therapy on catastrophising. The overall effect was just significant: $Z=1.99,\ P=0.05$ (Analysis 7.4). The effect size was SMD -0.72 (95% CI -1.43 to -0.01), but the I² value was 84%.

Behavioural versus treatment as usual at follow-up

We entered two studies with 182 participants into an analysis of the effects of behaviour therapy on pain at follow-up. The overall effect of behaviour therapy was not significant (Z = 0.21, P > 0.05) (Analysis 8.1). We entered three studies with 336 participants into an analysis of the effects of behaviour therapy on disability at follow-up: the overall effect of behaviour therapy was not significant (Z = 1.08, P > 0.05) (Analysis 8.2). We entered two studies with 160 participants into an analysis of the effects of behaviour therapy on mood at follow-up. The overall effect of behaviour therapy was not significant (Z = 0.90, P > 0.05) (Analysis 8.3). No studies provided data on catastrophising at follow-up for this comparison.

Pain outcomes

CBT appears to have a small effect on pain measured immediately post-treatment when compared with doing nothing (treatment as usual or waiting list), but not when compared with an active control, and there is no effect at follow-up. Behaviour therapy had no effect on pain compared to doing nothing, at either time point; there was only one study in the comparison with an active control, and that showed no benefit post-treatment or at follow-up.

Disability outcomes

CBT has a small effect on disability post-treatment and at followup, compared with an active control, and post-treatment compared with doing nothing, but this effect disappeared at followup. Behaviour therapy had no effect on disability compared to active control or to doing nothing, post-treatment or at followup, although there were only two studies comparing behaviour therapy with an active control.

Mood outcomes

CBT has no effect on mood immediately post-treatment compared with active control but, when compared with doing nothing (treatment as usual or waiting list), it has a moderate effect size immediately post-treatment and a small one at follow-up. Behaviour therapy had only one study in which it was compared with active control, and behaviour therapy showed no effect either post-treatment or at follow-up compared with doing nothing (treatment as usual or waiting list).

Catastrophising outcomes

CBT had a small effect compared to active control immediately post-treatment, lost at follow-up, but in comparison with doing nothing it had a moderate effect post-treatment which was sustained at follow-up. For behaviour therapy, study numbers were too small in the active control comparison, but in comparison with doing nothing, behaviour therapy had a small effect immediately post-treatment; there were no follow-up data.

Heterogeneity inspection

In the four analyses showing an effect of intervention over control but with high heterogeneity (I² > 50%), we undertook further exploratory analyses. By visual inspection we removed the outliers to test for their influence on the overall effect. In Analysis 3.2, heterogeneity was reduced to 55% by the removal of one positive outlier (Williams 1996), without affecting the overall significant result. In Analysis 4.2 and Analysis 4.3, removal of a single study (Van Koulil 2010) reduced heterogeneity to 0% but without changing the non-significant result (Analysis 4.2), and reduced it to 9% (Analysis 4.3) but also produced a non-significant result in place of the just significant one: Z was 1.57, P > 0.05. In Analysis 7.4, again removal of a single study (Thieme 2003) reduced heterogeneity to 0% and strengthened the result, although this was

now only produced by two studies: Z = 2.41 P < 0.05; the effect size was SMD -0.34 (95% CI -0.62 to -0.06).

Effects of quality ratings

We undertook three further analyses to assess the potential effects of quality. We excluded studies classified as 'high risk' for treatment quality from the analyses. This largely had the effect of increasing CBT effect sizes and reducing heterogeneity, but only in one case did it raise a small effect size to a moderate one: mood change for CBT compared to treatment as usual, post-treatment. There were no effects on behaviour therapy effect sizes where there were sufficient data to analyse. However, use of a compound quality rating scale such as Yates 2005 can be problematic (*Cochrane Handbook* chapter 8.3.3 (Higgins 2011)); although treatment quality was not associated with post-treatment N (rho = 0.17, P > 0.1), studies of higher quality were already well represented in effect sizes, and analyses are in any case weighted by sample size.

DISCUSSION

Evidence base

There is a large evidence base for estimating the effectiveness of psychological treatments in chronic pain. Before applying our new sample size criterion, we found 65 eligible trials, and these in turn came from a larger set which included trials whose psychological content or delivery was insufficient to convince us that the trial was of a genuine psychological treatment. All forms of psychological treatment were reviewed, ranging from well-established techniques such as biofeedback to more recent innovations such as acceptance and commitment therapy (Veehof 2011). Despite our strict criteria on psychological quality and size, we were able to use data from 35 randomised controlled trials (RCTs) (4788 treated participants) of specific behavioural or cognitive behavioural therapy. Cognitive behavioural therapy (CBT) and behaviour therapy dominate the evidence base; there were no trials of other psychological treatments such as psychodynamic or interpersonal psychotherapy, or dialectical behaviour therapy. We excluded the trials of mindfulness where they were based more on physical and meditative techniques than on cognitive or behavioural psychological techniques, and trials of specific methods which we judged fell outside CBT and which are already, at least in part, covered by other systematic reviews: internet intervention for pain (Bender 2011; Macea 2010) and self regulation (in rheumatoid arthritis: Knittle 2010). A systematic review of hypnosis is due given its resurgence as a treatment method (Jensen 2011), albeit aimed more at pain reduction than overall rehabilitation.

This review includes 22 trials from the previous systematic review and 20 new or re-entered trials; seven trials provided no useable

data. The remainder allowed for reasonable power in the analyses, with the largest analysis being of 1258 participants (CBT versus active control post-treatment) and the smallest of 144 participants (behaviour therapy versus active control at follow-up). An analysis of quality scores, as measured by the Yates et al scale (Yates 2005), showed that the quality of the design and reporting of trials has clearly improved over the years, perhaps as a consequence of the emphasis of Cochrane and other evidence-focused organisations concerned with methodological standards such as CONSORT (Boutron 2008). However, the quality of treatments, of their reporting, or both, does not appear to have improved over time, but the Yates 2005 treatment subscale is restricted to five items so may be relatively insensitive.

Summary of results

The majority of studies were of CBT, reflecting its dominance in chronic pain management and in psychological treatment more widely. Of the eight comparisons of CBT versus active control (four outcomes: pain, disability, mood, catastrophic thinking, at two time points, immediately post-treatment and six to 12 months follow-up), three were positive: disability immediately post-treatment and at follow-up, and catastrophic thinking post-treatment. There were stronger effects for the seven comparisons of CBT versus doing nothing (treatment as usual or waiting list) for four outcomes: small effects on pain and on disability post-treatment but not at follow-up; small effects on mood maintained at follow-up; and moderate effects on catastrophising with insufficient data to analyse at follow-up.

For behaviour therapy the evidence is much weaker and, with our more stringent criteria, rather sparse. Behaviour therapy was developed in the 1960s and 1970s and evaluated as part of the first wave of psychological treatment for pain (Morley 2011). As a consequence, trials tended to be small and methodologically weak and have been largely superseded by procedures that claim to be cognitive behavioural. Compared with doing nothing (treatment as usual or waiting list), behaviour therapy has no effects on pain, disability or mood immediately post-treatment, but a small effect on catastrophic thinking; there were insufficient data at follow-up except for disability, where there was no effect.

The size of effects - small to moderate - is similar to other systematic reviews in this field: of mixed chronic pain (Scascighini 2008), low back pain (Henschke 2010a; Hoffman 2007), fibromyalgia (Bernady 2010; Glombiewski 2010; Häuser 2009) and arthritis (Dixon 2007). It is also comparable with effect sizes of CBT for pain problems in children (Eccleston 2009b) and for major psychological disorders (Butler 2006). Of our four outcome domains, effects on mood (mostly depression) were strongest, followed by catastrophic thinking, disability and, lastly, pain. We did not include reduction in health care use (but see Bernady 2010), or cost-effectiveness (see Gatchel 2006).

Change in evidence from previous review

We raised the quality criterion for N (sample size) in this review, bearing in mind the risk of bias of small numbers in trials (Nuesch 2009), and the overall tendency for poorer quality trials to produce more positive results (Furlan 2001; Ioannidis 2005; Nuesch 2009). Numbers in trials have steadily increased over time, but in some cases this appears to be at the cost of treatment intensity (for instance, number of hours of patient contact, or staff experience). Compared to our 2009 review (Eccleston 2009a), the effect sizes for CBT are largely sustained and extended with the addition of catastrophic thinking as an outcome, while those for behaviour therapy are diminished. Treatment gains are of the same order as those of other available treatments (Glombiewski 2010), as shown by head-to-head trials of surgery versus psychologically based rehabilitation (Fairbank 2005; Hellum 2011).

Issues for consideration

Psychological therapies for the management of chronic pain are potentially useful treatments, with better evidence for and better effects of CBT than behaviour therapy. There are, however, many problems in interpreting the data and using it to devise a strategy to improve our understanding. We discuss the most important issues, or those that particularly affect psychological treatments, below, and then discuss what we should do next as a research community instead of simply continuing to conduct small RCTs and systematically reviewing them.

- 1. The lack of coherent theory underlying many of these studies remains a concern. We do not have a clear notion of the mechanisms of change in CBT trials (pace Jensen 2011), nor are we yet able to distinguish well between specific effects of therapy and nonspecific effects of the interactions and context, an unresolved issue in psychology more broadly (Roth 2005). A simple model of independent deficits in cognition, emotion or physical function to be remedied by independent components of therapy is inadequate; even assumptions of deconditioning and poor physical status in chronic pain have proved to be unsubstantiated (Lin 2011; Verbunt 2010). Change in some outcomes may be needed to facilitate change in others: it is common to assume or to hypothesise that change in beliefs and ways of thinking (such as catastrophising) mediates other changes (Moss-Morris 2007; Thorn Burns 2011), and this will not be tested by RCTs or post hoc data analyses (even in very large trials: Underwood 2011) but by carefully designed prospective studies (Wideman 2009) and experimental analysis of specific treatment components (e.g. Vlaeyen de Jong
- 2. There are particular issues of bias and potential bias in trials which affect interpretation of results and conclusions to be drawn from them. These can be described under the headings of patient factors, treatment factors and methodological issues.
- 2.1. Patient samples are heterogeneous (Turk Okifuji 2002) and,

without a suitable theory, our attempts to subgroup them are either based on non-psychological properties, such as diagnosis, or on superficial, non-functional characteristics which can be elicited by questionnaire. Neither strategy is likely to be helpful in identifying what works for whom. A more psychologically informed subgrouping of patients, rather than by diagnostic group, should allow better targeted and more effective treatment (Morley 2006). Matching patients to treatment components according to baseline problem severity misses the demonstrated impact of, for instance, the behavioural component on emotional problems, or the cognitive component on physical activity. Treatment makes substantial demands on patients, although many trials do not monitor whether patients practise treatment components as instructed. Treatment aims to enable long-term changes in behaviour related to pain, but a test of adherence in the month following intensive CBT to cognitive, exercise and activity plans showed only 2% to 3% of variance in outcomes explained by adherence (Curran 2009). While assessment of adherence could doubtless be improved, we strongly suspect that the model of adherence is too simple, failing to look beyond the patient and to acknowledge substantial obstacles wholly or partly outside the patient's control (Nicholas 2010).

2.2. Treatments are similarly heterogenous, and the procedures included in treatment arms of many of the trials reviewed are pragmatic mixes of various content, often without an adequate rationale, and with apparent disjunction between stated aims of treatment, actual treatment content and outcomes measured. Component dismantling studies offer an illusion of identifying 'active ingredients' of the total package when we do not yet have the power of numbers, nor the statistics, to calculate the effects of each component on each outcome (Grimshaw 1995). Treatment content is difficult to represent even given the possibility of extended accounts on internet appendices (Thorn 2007); and, although it is possible to measure treatment fidelity (Leeuw Goossens 2008), we still do not know whether the unique components of therapy are the important ones. The heterogeneity indices give reason to suspect that there are important differences between treatments of potential interest that have yet to be identified. Treatment content, even with the most detailed protocol, will differ in the hands of different therapists with greater or lesser skills at eliciting and working with examples of emotional and practical importance for patients rather than talking in general terms about change, a particular issue of concern with less experienced therapists (Waller 2009). Then any psychological treatment has to be, as it were, manufactured in the moment it is delivered. In this way psychological therapies are comparable with surgery, rather than with pharmacotherapy.

2.3. There are further particular methodological issues of note. In particular, patients presenting with multiple problems captured by the label of chronic pain, treated with multicomponent, often programmatic, treatments, unsurprisingly make many changes which may or may not be captured by outcome measures. Neither pa-

tients nor trial authors agree on the relative importance of all targets of treatment (Beale 2011; Turk 2008). Further, outcomes are analysed as if independent, although they are unlikely to be so. In a few cases this has been empirically demonstrated: depression and physical disability tend to be associated in chronic pain, independent of measurement contamination (Alschuler 2008). While standard reporting (Brown Brunnhuber 2006; Garratt 2008; Thorn 2007) would help to some extent, the problem lies in the lack of adequate models to guide intervention. The field should seriously consider developing measures which are capable of indexing clinical improvement to replace or augment statistical change (Morley 2006) and which have ecological validity. Broad spectrum measures of the disability domain, such as quality of life (e.g. Short Form 36 Health Survey) may have validity problems when applied to trials of the effectiveness of therapies, caused largely by the inclusion of content either irrelevant to the patient, not the target of treatment, or both (Bowling 1997; Dworkin 2005). As a consequence, the sensitivity of measures may be compromised. Additionally, trials report results in terms of statistical rather than clinical significance, which may have led to earlier optimistic summaries of effectiveness. Binary outcomes based on a clinical significance criteria (Morley 2006; Morley 2008) would allow us to estimate treatment responders (Dworkin 2005; Dworkin 2008): people who are 'successfully' treated by CBT or behaviour therapy, and to estimate adverse events, the lack of attention to which is deplorable. We note that a recent study of effectiveness observed evidence of deterioration in a small proportion of patients using statistically defined criteria for clinically significant change (Morley 2008). 3. A further methodological issue is the design of control groups. Relatively few trials in this review used 'attention control' structurally equivalent to the active treatment, with the explicit aim of minimising differences between conditions in such nonspecific effects, and our separation of comparisons of active comparators from treatment as usual or waiting list does not match the category of attention control. Particularly in studies that compare mean data from continuous measures (Hrobaritsson 2001; Hrobaritsson 2004), this leaves uncertainty about whether the benefits of treatment can be attributed to specific features of treatment. This is not unique to psychological studies but it is relatively rare for it to be acknowledged in studies of physical or drug interventions (Wren 2011, yoga review). We strongly suspect that as a field we have underestimated the complexity of behaviour change and the social and psychological influences that maintain disability in chronic pain patients (Blyth 2007). Further, the typical chronic pain patient has well-established behavioural patterns reinforced over a long period of failed attempted adjustment to pain and distress, and it has not been established whether the psychotherapeutic content of existing trials is adequate; the current review cannot resolve that question. While it is possible, and we plan, to perform sub-analyses for various aspects of treatment, such as treatment intensity or quantity (associated with outcome by several other systematic reviews: Scascighini 2008, Glombiewski 2010 and in psychological treatments in general, e.g. Barkham 2006), those aspects of treatment such as treatment content, quantity (dose), staff competence and patient population are not independent of each other in their effects on outcome. We speculate, however, that good clinical outcomes should perhaps not be expected from dilute and brief treatments delivered by inexperienced staff to severely distressed patients, particularly given the poor preparation and access to specialist care identified in primary care studies (Breivik 2006; Gatchel 2006; Somerville 2008).

4. We know that the effects of drugs in chronic pain tend to be either very good or very poor. Response is bimodal, with small numbers of responders, often as little as 5% to 20% more than with placebo drug (Moore 2010). It may well be that psychological intervention has a similar type of response, with a small number of patients making substantial changes but most changing little, making trials and meta-analyses relatively insensitive. This review has used average scores because average scores are reported in trials. It is arguably more relevant to analyse data by the number of individual patients achieving a level of longer-term improvement in pain, disability, distress or other problem; a level set with reference to clinical meaning. In some chronic pain conditions which are difficult to treat, like fibromyalgia or chronic low back pain, the proportion of patients benefiting from drug treatment is small. Similar low success rates are likely for psychological interventions, especially in populations in which many previous treatments have failed.

AUTHORS' CONCLUSIONS

Implications for practice

Psychological interventions can reduce pain, disability, psychological distress and catastrophic ways of thinking about pain. Average effect sizes derived from collapsing data across trials are relatively small, as they are across pharmacological and physical treatments for chronic pain. Examination of what we think is feasible as the outcome of psychological treatment is appropriate: is it mere palliation, in which case effects will be small, or do we expect to move people who are stuck in trying to solve the unsolvable problem of pain to address instead the solvable problem of living more satisfactorily with chronic pain (Eccleston 2007), and starting to do so? Or to put it another way, do we believe that we effectively enable patients to manage the interruption of pain and to reduce its interference with their lives, and thereby to repair damaged identities (Morley 2011). These are substantial changes, unlikely to occur rapidly (within the timescale of some trials). What is evident from this review is the following:

1. CBT is effective when delivered by experienced staff, those trained and supervised in the trial protocol, or both. The results cannot be extrapolated to CBT delivered by untrained staff.

- 2. There is no clear benefit of adding further components to multicomponent CBT: it is unlikely that the extra component, such as two sessions on 'mindfulness', will make any measurable difference. The rationale given for such additions in trials in this review was often weak.
- 3. Although trials do not tend to report adverse effects or deterioration (such as worsening of depression to a level of clinical concern), we know that such effects should be small (Fairbank 2005; Hellum 2011; Morley 2008), so the treatment can be considered to be safe, with the reservation that the reasons for discontinuing treatment are rarely given and may be due to hidden adverse effects.
- 4. Average effects mask larger changes on the part of some patients and little or none for others. Better trial design and observational studies will help us to identify those patients for whom CBT can enable substantially better outcomes, and those who need current treatment to be adapted or who need other treatment to improve their quality of life with chronic pain. Clinicians can contribute significantly to generating hypotheses about how to distinguish these patients from one another.
- 5. The way forward for psychological treatment lies not in more RCTs, unless the intervention is entirely novel, the patient population has not previously been studied, or the outcomes are truly innovative. Any new RCT needs to be designed and reported taking explicit account of the challenges identified and discussed in this review.

Implications for research

- 1. We recommend the immediate cessation of new RCTs of CBT against simple alternatives, unless a strong case can be given for the novelty of the population or treatment under investigation. We include in this recommendation treatments of CBT with additional components: see Implications for practice, point 2. The evidence of weak to moderate effects across a range of outcomes is clear from our systematic reviews and from the others cited above, and is very unlikely to change as a result of further similar RCTs and systematic reviews. The average effects are small, as they are for all treatments of chronic pain (Moore 2010).
- 2. The question addressed by psychological treatment for chronic pain is complex, conceptually and statistically. We no longer believe that it is possible to design a 'pure' trial of a single component of intervention (such as relaxation, operant reinforcement or acceptance), although that is not to deny that there is much to be learned from some of the trials which attempt it in this review. Suggested solutions in realistic and clinically informed evaluations of complex packages (Craig 2008; Shepperd 2009) will take us no further than the current review and, with pressure to economise on resources, there is so far no indication of which components should be cut or retained.

- 3. Since we share these challenges with the larger field of pain medicine, we can usefully consider some current initiatives: running N of 1 trials (McMillan 2010); examining individual data for response trajectories (Lambert 2001; Moore 2005); pooling data for responder analyses (Moore 2010); or conducting clinical effectiveness trials (Moore 2010), where 'clinical effectiveness' is "the product of efficacy, tolerability, utility, cost, and speed" (Moore 2010, p174) so that trials focus on maximising benefit and minimising cost, including adverse events.
- 4. We need better theory to generate hypotheses about processes and mechanisms of change, to be tested in terms of

populations, treatment content, treatment process and outcomes.

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^{*} Indicates the major publication for the study

CHARACTERISTICS OF STUDIES

Characteristics of included studies [ordered by study ID]

Alaranta 1994

Methods	RCT; 2 arms;assessed at pretreatment, 3 months follow-up, 1 year follow-up
Participants	3 month follow-up n = 286 Start of treatment n = 293 Sex: 160 F, 133 M Mean age = 40.5 (SD 4.5) Source = patients referred for inpatient rehabilitation Diagnosis = chronic low back pain Mean years of pain = not given (minimum 6 months)
Interventions	"progressive intervention of intensive physical training and psychosocial activation AK- SELI" "control: less intensive physical training and passive physical therapies"
Outcomes	Primary pain outcome: none Primary disability outcome: none
	Primary mood outcome: BDI
	Catastrophising outcome: none
	1. lumbar flexion-extension
	2. lateral flexion
	3. trunk rotation
	4. hamstring tightness
	5. number of sit-ups
	6. number of arch-ups
	7. static strength of back muscles
	8. number of squats
	9. Million index of pain and disability mean of 14 items rated 0 to 100
	10. low back pain capacity 1 to 3
	11. leisure activities physical intensity 0 to 10
	12. number of visits to doctors (12-month follow-up)
	13. number of physical therapy outpatient visits (12-month follow-up)
	14. WHO occupational handicap 0 to 5
	15. sick days
	16. Beck Depression Inventory
	17. Symptom Check List
	18. Multidimensional Health Locus of Control
	19. Social Adjustment Scale
	20. Karolinska Scales of Personality
Notes	Excluded from 2009 review for marginal psychological content; included in 2012 update No data Yates quality scale: total quality = 16/35, design quality = 13/26, treatment quality = 3/
	9

Alaranta 1994 (Continued)

Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	"patients stratified according to age and sex and randomly divided into intervention and control groups"
Allocation concealment (selection bias)	High risk	No information but post-randomisation exclusion of patients "not fit" for intervention group
Incomplete outcome data (attrition bias) All outcomes	High risk	Attrition implied not reported; no reporting of differences
Selective reporting (reporting bias)	High risk	Many outcomes not reported
Blinding of outcome assessment (detection bias) All outcomes	High risk	Self report and examination by physiatrist and physiotherapist at baseline and follow- up. No statement about blinding

Altmaier 1992

Methods	RCT; 2 arms; assessed at pre-treatment, post-treatment, 6 months
Participants	End of treatment n = 42 Start of treatment n = 45 Sex: 12 F, 33 M Mean age = 39.9 (SD 8.9) Source = pain and rehabilitation clinic Diagnosis = chronic low back pain Mean years of pain = not given
Interventions	"Psychology based programme: multicomponent CBT" "Standard inpatient rehabilitation"
Outcomes	Primary pain outcome: MPQ PRI Primary disability outcome: WHYMPI pain interference Primary mood outcome: WHYMPI distress Catastrophising outcome: none 1. Primary aerobic impairment 2. Self efficacy 3. West Haven Yale Multidimensional Pain Inventory (WHYMPI) self control 4. West Haven Yale Multidimensional Pain Inventory (WHYMPI) pain interference 5. West Haven Yale Multidimensional Pain Inventory (WHYMPI) mood 6. Disability 7. Melzack Pain Questionnaire Pain Response Index (MPQ PRI)

Altmaier 1992 (Continued)

Notes	CBT versus TAU, post-treatment and follow-up: analyses 3.1, 3.2, 3,3, 4.1, 4.2, 4.3
	Yates quality scale: total quality = 15/35, design quality = 11/26, treatment quality = 4/
	9

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	Abstract: "Forty-five low back pain patients were randomly assigned"; no details in Methods
Allocation concealment (selection bias)	High risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	High risk	Inadequately reported
Selective reporting (reporting bias)	Low risk	Fully reported
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not reported

Basler 1997

Methods	RCT; 2 arms; assessed at pre-treatment, post-treatment, 6 months
Participants	End of treatment n = 76 Start of treatment n = 94 Sex: 57 F, 19 M Mean age = 49.3 (SD 9.7) Source = pain or rehabilitation clinic Diagnosis = chronic low back pain Mean years of pain = 10.8
Interventions	"CBT added to medical treatment" "Medical treatment"
Outcomes	Primary pain outcome: NRS 0 to 10 pain Primary disability outcome: disability in physical function from Dusseldorf Disability Scale Primary mood outcome: none Catastrophising outcome: PRSS catastrophising 1. Pain Intensity Numerical Rating Scale (0 to 10) 2. Control over pain Numerical Rating Scale (0 to 10) 3. Days per week pain-free 4. Days per week pain medication use 5. Use of cognitive strategies (self report)

Basler 1997 (Continued)

	 6. Use of avoidance behaviour (self report) 7. Pleasant activities (self report) 8. Social support (self report) 9. Philosophical beliefs (self report) 10. Catastrophising (bespoke scale) 11. Active coping (bespoke scale) 12. Disability in social relationships from Dusseldorf Disability Scale 13. Disability in social roles from Dusseldorf Disability Scale 14. Disability in physical function from Dusseldorf Disability Scale 15. Disability in mental performance from Dusseldorf Disability Scale
	15. Disability in mental performance from Dusseldorf Disability Scale 16. Disability in physical performance from Dusseldorf Disability Scale
Notes	CBT versus TAU, post-treatment: analyses 3.1, 3.2 Yates quality scale: total quality = 18/35, design quality = 12/26, treatment quality = 6/9

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	"Through assignment of random numbers, patients were allocated to an experimental treatment or a control group."
Allocation concealment (selection bias)	High risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition reported; 1 difference found between dropouts and completers
Selective reporting (reporting bias)	Low risk	Fully reported
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not reported

Bliokas 2007

Methods	RCT; 3 arms; assessed at pretreatment and post-treatment
Participants	End of treatment n = 94 Start of treatment n = 143 Sex: 79 F, 64 M Mean age = 45.2 (SD 9.2) Source = referrals to Pain Management Service after medical treatment completed Diagnosis = chronic non-cancer pain Mean years of pain = median 4.0

Bliokas 2007 (Continued)

Interventions	"Graded exposure in vivo and outpatient multidisciplinary chronic pain management group program" "outpatient multidisciplinary chronic pain management group program" "Waiting list control"
Outcomes	Primary pain outcome: pain VAS Primary disability outcome: Pain Disability Index Primary mood outcome: DASS depression Catastrophising outcome: none 1. Pain VAS 2. Tampa Scale for Kinesiophobia: fear of movement/re/injury 3. Pain Self-Efficacy Questionnaire (PSEQ) 4. Pain Disability Index (PDI) 5. Depression, Anxiety & Stress Scale (DASS): depression and anxiety scores only 6. Activity level: performance over 2 weeks of 10 usually avoided activities 7. 6-minute walk test
Notes	Chronic pain management programme with graded exposure versus waiting list control December 2009 search Data obtained from author: analyses 3.1, 3.2, 3.3 Yates quality scale: total quality = 23/35, design quality = 15/26, treatment quality = 8/9

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Random numbers generation
Allocation concealment (selection bias)	High risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition fully reported; no differences
Selective reporting (reporting bias)	Low risk	Fully reported
Blinding of outcome assessment (detection bias) All outcomes	High risk	Examination by physiotherapist and self report: no blinding reported

Buckelew 1998

Methods	RCT; 4 arms; assessed at pre-treatment, post-treatment, 3 months, 1 year, 2 years
Participants	End of treatment n = 109 Start of treatment n = 119 Sex: 108 F, 11 M

Buckelew 1998 (Continued)

	Mean age = 44 (SD 10) Source = mainly community Diagnosis = fibromyalgia Mean years of pain = 11.5
Interventions	"Biofeedback + relaxation + exercise" "Biofeedback + relaxation" "Exercise" "Education attentional control"
Outcomes	Primary pain outcome: no data available Primary disability outcome: no data available Primary mood outcome: no data available Catastrophising outcome: no data available Arthritis Impact Measurement Scale: Physical Activity subscale (AIMS) Symptom Checklist (SCL-90R) distress Center for Epidemiologic Studies Depression Scale (CES-D) Arthritis Self-Efficacy Scale Sleep rating 0 to 12 Tender Point Index Myalgic score Physician's VAS rating of disease severity Keefe & Block Pain Behaviour: observation
Notes	No data Yates quality scale: total quality = 20/35, design quality = 15/26, treatment quality = 5/9

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	"randomly assigned"
Allocation concealment (selection bias)	High risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Attrition partially reported and did not differ across groups; no test for differences
Selective reporting (reporting bias)	Low risk	Fully reported
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Subjects examined by physician unaware of treatment conditions and with no other contact with subjects

De Souza 2008

Methods	RCT; 2 arms; assessed at pre-treatment, post-treatment, 4 months, 12 months
Participants	End of treatment n = 55 Start of treatment n = 60 Sex: 60 F, 0 M Mean age = 49.6 (SD 7.0) Source = not stated Diagnosis = fibromyalgia Mean years of pain = 12.4
Interventions	"Interactional School of Fibromyalgia" "Control"
Outcomes	Primary pain outcome: MPI pain severity Primary disability outcome: MPI interference with daily activity Primary mood outcome: none Catastrophising outcome: none 1. VAS pain (pain diary) 2. MPI pain severity 3. MPI pain interference daily activity 4. MPI control over pain 5. MPI mood 6. MPI family and social support 7. VAS suffering (pain diary) 8. VAS ability to do daily activity (pain diary)
Notes	December 2009 search No data Yates quality scale: total quality = 13/35, design quality = 10/26, treatment quality = 3/9

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	"randomly assigned"
Allocation concealment (selection bias)	High risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	High risk	Attrition partially reported; no test for differences
Selective reporting (reporting bias)	Low risk	Fully reported
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not reported

Ehrenborg 2010

Methods	RCT; 2 arms; assessed at post-treatment and 6-month follow-up
Participants	End of treatment: n = 62 Start of treatment: n = 65 Sex: 33 F, 29 M Age: mean = 39.4 (SD 11.1) Mean years of pain = 2.1 (SD 2.5) Source = outpatient rehabilitation unit Diagnosis = pain (neck and shoulder) after whiplash injury
Interventions	CBT rehabilitation plus EMG biofeedback versus CBT rehabilitation
Outcomes	Primary pain outcome: no data Primary disability outcome: Canadian Occupational Performance Measure Primary mood outcome: none Catastrophising outcome: none Canadian Occupational Performance Measure Multi-dimensional Pain Inventory (Swedish)
Notes	CBT versus active, post-treatment and follow-up: analyses 1.2 and 2.2 2011 search Yates quality scale: total quality = 21/35, design quality = 17/26, treatment quality = 4/9

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"Randomization was performed by casting a die after the participant's acceptance: odd numbers for treatment group"
Allocation concealment (selection bias)	High risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition fully reported
Selective reporting (reporting bias)	Low risk	Fully reported
Blinding of outcome assessment (detection bias) All outcomes	High risk	Therapists conducted assessments: statement that study not blinded

Ersek 2008

Methods	RCT; 2 arms; assessed at pretreatment, post-treatment, 6-month follow-up, 12-month follow-up
Participants	End of treatment n = 218 Start of treatment n = 256 Sex: 210 F, 46 M Mean age = 81.8 (SD 6.5) Source = residential retirement facilities Diagnosis = pain more than 3 months; average last week > 2/10: mixed sites, largest legs and feet Mean years of pain = not given
Interventions	"pain self-management training group (SMG) intervention" "education only control condition"
Outcomes	Primary pain outcome: BPI pain Primary disability outcome: RMDQ Primary mood outcome: Geriatric Depression Scale Catastrophising outcome: CSQ catastrophising 1. Roland & Morris Disability Questionnaire 2. Brief Pain Inventory: pain 3. Brief Pain Inventory: interference with activity 4. Geriatric Depression Scale 5. Arthritis Self-Efficacy Scale 6. CSQ catastrophising 7. Chronic Pain Coping Inventory: guarding 8. Chronic Pain Coping Inventory: resting 9. Chronic Pain Coping Inventory: relaxation 11. Chronic Pain Coping Inventory: task persistence 12. Chronic Pain Coping Inventory: exercise/stretch 13. Chronic Pain Coping Inventory: seeking support 14. Chronic Pain Coping Inventory: coping self statements 15. Chronic Pain Coping Inventory: pacing 16. Medication use: record
Notes	December 2009 search: 1.1, 1.2, 1.3, 1.4, 2.1, 2.2, 2.3, 2.4 Yates quality scale: total quality = 30/35, design quality = 21/26, treatment quality = 9/9

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomisation by (retirement) facility, by statistician using random number generator
Allocation concealment (selection bias)	Low risk	By independent statistician

Ersek 2008 (Continued)

Incomplete outcome data (attrition bias) All outcomes	Low risk	Fully reported attrition
Selective reporting (reporting bias)	Low risk	Fully reported
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not reported

Evers 2002

Evers 2002	
Methods	RCT; 2 arms; assessed at pre-treatment, post-treatment, 6 months follow-up
Participants	End of treatment n = 59 Start of treatment n = 64 Sex: 42 F, 17 M Mean age = 54.1 (SD 11.4) Source = rheumatology clinic Diagnosis = rheumatoid arthritis Mean years of pain = 3.1
Interventions	"Tailor made CBT" "Treatment as usual"
Outcomes	Primary pain outcome: IRGL Pain Primary disability outcome: IRGL Functional Disability (Composite Z score) Primary mood outcome: BDI depression Catastrophising outcome: Illness Cognitions - Helplessness Disease Activity Invloed van Reuma op Gezondheid en Leefwijze (IRGL): Functional Disability Invloed van Reuma op Gezondheid en Leefwijze (IRGL): Pain Invloed van Reuma op Gezondheid en Leefwijze (IRGL): Anxiety Invloed van Reuma op Gezondheid en Leefwijze (IRGL): Perceived support Social network Illness Cognitions: Helplessness Illness Cognitions: Acceptance Active Coping with Pain Passive Coping with pain Active Coping with Stress Passive Coping with Stress Passive Moping with Stress Fatigue Beck Depression Inventory Negative Mood (ZwartSpooren) Medication compliance
Notes	CBT versus TAU, post-treatment and follow-up: analyses 3.1, 3.2, 3.3, 4.1, 4.2, 4.3 Yates quality scale: total quality = 25/35, design quality = 18/26, treatment quality = 7/9

Evers 2002 (Continued)

Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Random numbers
Allocation concealment (selection bias)	Low risk	"previously determined"
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition fully reported
Selective reporting (reporting bias)	Low risk	Fully reported
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not reported

Falcao 2008

Methods	RCT; 2 arms; assessed at pre-treatment, post-treatment, 3 months
Participants	End of treatment n = 51
	Start of treatment $n = 60$
	Sex: 60 F, 0 M
	Mean age = 45.7 (SD 2.3)
	Source = Rheumatology outpatients
	Diagnosis = fibromyalgia
	Mean years of pain = 3.6
Interventions	"Cognitive behavioral therapy"
	"Routine medical visits"
Outcomes	Primary pain outcome: VAS
	Primary disability outcome: FIQ (no data for SF-36)
	Primary mood outcome: BDI
	Catastrophising outcome: none
	1. Visual analogue scale for pain
	2. Verbal improvement scale (5 categories)
	3. Fibromyalgia Impact Questionnaire (FIQ)
	4. SF-36 physical capacity (function)
	5. SF-36 physical aspects (role)
	6. SF-36 pain
	7. SF-36 general health
	8. SF-36 vitality
	9. SF-36 social aspects
	10. SF-36 emotional aspects
	11. SF-36 mental health

Falcao 2008 (Continued)

	12. Beck Depression Inventory (BDI)13. Spielberger State-Trait Anxiety Inventory (STAI State)14. Number of paracetamol tablets
Notes	December 2009 search: analyses 3.1, 3.2, 3.3 Yates quality scale: total quality = 16/35, design quality = 14/26, treatment quality = 2/9

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"Patients were randomized by drawing lots"
Allocation concealment (selection bias)	High risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Attrition partially reported; statement that dropouts were not different
Selective reporting (reporting bias)	Low risk	Fully reported
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Evaluation by clinician blind to treatment allocation

Geraets 2005

Methods	RCT; 2 arms; assessed at pre-treatment, post-treatment, 1 year
Participants	End of treatment n = 158 Start of treatment n = 176 Sex: 109 F, 83 M (at start of treatment) Mean age = 52.5 (SD 12.4) Source = mixed community and volunteer Diagnosis = shoulder pain Mean years of pain = not given
Interventions	"Graded exercise" "Primary care TAU"
Outcomes	Primary pain outcome: NRS Primary disability outcome: Shoulder Disability Questionnaire Primary mood outcome: none Catastrophising outcomes: PCCL catastrophising Shoulder disability questionnaire Shoulder pain Pain intensity NRS

Geraets 2005 (Continued)

	Quality of life
	Fear avoidance
	Kinesiophobia (2 items)
	Pain Coping and Cognition List: catastrophising
	Pain Coping and Cognition List: coping
	General Practitioner visits
	Physician visits
	Physiotherapy visits
	Number of drug prescriptions
	Number of days work absence
	Total cost of health care (Euros)
Notes	BT versus TAU: analyses 7.1, 7.2, 7.4, 8.2 Yates quality scale: total quality = 26/35, design quality = 20/26, treatment quality = 6/9

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Block randomisation according to random number table
Allocation concealment (selection bias)	Low risk	Random number table generated by person not involved in study; opaque sealed envelopes; "Blinding for patients of allocated treatment was not possible" but treatment preferences elicited and shown to have no effect on outcome
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition fully reported; dropouts different in pain characteristics but not outcome measures at baseline
Selective reporting (reporting bias)	Low risk	Fully reported
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Researchers not involved in randomisation collect data

Glombiewski 2010b

Methods	RCT; 3 arms: CBT + biofeedback; CBT; waiting list control, post-treatment (WLC assigned to treatment so no WLC at 6-month follow-up)
Participants	End of treatment: n = 116 Start of treatment: n = 128 Sex: 77 F, 39 M Mean age: 48.8 (SD 11.7)

Glombiewski 2010b (Continued)

	Source = medical referrals (86%) or response to newspaper advert (14%) Diagnosis = chronic back pain Mean years of pain: 8.1 (SD 8.7)
Interventions	"CBT with biofeedback" "CBT" "waiting list control"
Outcomes	Primary pain outcome: 0 to 10 NRS pain intensity Primary disability outcome: PDI Primary mood outcome: BDI Catastrophising outcome: none Pain intensity 0 to 10 NRS Pain average of 4x daily diary for 1 week Pain Disability Index Beck Depression Inventory Coping Strategies Scale from FESV Health-Related Life Satisfaction Scale Global treatment change Treatment satisfaction (Adverse events noted from pain intensity and global treatment change) Health care use: doctor visits for pain
Notes	Combined (CBT + biofeedback and CBT) versus WLC: analyses 3.1, 3.2, 3.3 2011 update search Yates quality scale: total quality 24/35, design quality 17/26, design quality 15/26

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomisation by random number generation
Allocation concealment (selection bias)	Unclear risk	"coordinated by the first author" before study
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Attrition fully reported; statement that dropout data will be reported elsewhere
Selective reporting (reporting bias)	Low risk	Fully reported
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not reported

Greco 2004

Methods	RCT; 3 arms; assessed pre-treatment, post-treatment, 6/9 months
Participants	End of treatment n = 80 Start of treatment n = 92 Sex: 87 F, 5 M (at start of treatment) Mean age = 47.3 (SD 10.4) Source = volunteers Diagnosis = SLE Mean years of pain = 11
Interventions	"CBT with biofeedback" "Symptom monitoring and support" "Treatment as usual"
Outcomes	Primary pain outcome: AIMS2 pain 0 to 10 Primary disability outcome: SF36 physical function (reversed) Primary mood outcome: CES-D Depression Catastrophising outcome: perceived stress Arthritis Impact Measurement Scale (AIMS) 2: pain Multidimensional Pain Inventory: interference Center for Epidemiologic Studies Depression Scale (CES-D) Arthritis Self-Efficacy Perceived stress Short Form 36 physical function Fatigue severity Global self assessment Disease activity systemic lupus activity measure-revised (SLAM-R) Systemic Lupus Erythematosus Disease Activity Index (SLEDAI)
Notes	CBT versus active, post-treatment and follow-up: analyses 1.1, 1.2, 1.3, 2.1, 2.2, 2.3 CBT versus TAU, post-treatment and follow-up: analyses 3.1, 3.2, 3.3, 4.1, 4.2, 4.3 Yates quality scale: total quality = 25/35, design quality = 20/26, treatment quality = 5/9

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	"assigned randomly, based on a software- generated randomization plan"
Allocation concealment (selection bias)	High risk	Not reported, but equal credibility of treatments rated by participants
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Attrition fully reported; no test for differences
Selective reporting (reporting bias)	Low risk	Fully reported

Greco 2004 (Continued)

Blinding of outcome assessment (detection bias)	Low risk	Rheumatologist and researcher assessors masked to treatment assignment
All outcomes		Ü

Haldorsen 1998

Methods	RCT; 2 arms; assessed pre-treatment, 1 year
Participants	End of treatment n = 387
•	Start of treatment $n = 469$
	Sex: 298 F, 171 M
	Mean age = 43 (SD 10.6)
	Source = National Insurance system contact
	Diagnosis = mixed chronic pain
	Mean years of pain = not given
Interventions	"Cognitive behaviour therapy"
	"Treatment as usual"
Outcomes	Primary pain outcome: VAS pain
Primary disability outcome: none	
	Primary mood outcome: HSCL distress
	Catastrophising outcome: none
	Visual analogue scale pain (in afternoon)
	Physical training
	Hopkins Checklist (HSCL) Distress (Norwegian version)
	Attribution style Work satisfaction
	Ergonomic performance
	Subjective health rating
Notes	CBT versus TAU post-treatment: analyses 4.1, 4.3
	Yates quality scale: total quality = 12/35, design quality = 10/26, treatment quality = 2/
	9

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Allocated at random by cards in sealed envelopes
Allocation concealment (selection bias)	Low risk	Allocation sequence by someone not involved in study
Incomplete outcome data (attrition bias) All outcomes	High risk	Attrition partially reported; no test for differences

Haldorsen 1998 (Continued)

Selective reporting (reporting bias)	High risk	Not fully reported
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessment by physiotherapists who tried to remain blind to treatment

Hammond 2001

Methods	RCT; 2 arms; assessed at pre-treatment, post-treatment, 1 year
Participants	End of treatment n = 121 Start of treatment n = 127 Sex: 97 F, 30 M Mean age = 50.5 (SD 10.6) Source = rheumatology clinic Diagnosis = rheumatoid arthritis (hand) Mean years of pain = 1.6
Interventions	"Joint protection arthritis education" "Standard arthritis education"
Outcomes	Primary pain outcome: none available Primary disability outcome: AIMS2 activities of daily living Primary mood outcome: none available Catastrophising outcome: RAI helplessness Adherence to joint protection Hand pain visual analogue scale Overall pain visual analogue scale Tender count (28 joints) Swollen joint count (28 joints) Early morning stiffness Grip strength Hand joint alignment Arthritis Impact Measurement Scale (AIMS) 2: ADL Arthritis Impact Measurement Scale (AIMS) 2: lower limb function Arthritis Impact Measurement Scale (AIMS) 2: current health status Arthritis Self Efficacy (pain) Arthritis Self Efficacy (other) Rheumatoid attitude index (helplessness) Rheumatoid attitude index (internality) Satisfaction with health
Notes	CBT versus active, post-treatment and follow-up: analyses 1.2, 2.2 Yates quality scale: total quality = 18/35, design quality = 15/26, treatment quality = 3/9

Hammond 2001 (Continued)

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	"allocated randomly"
Allocation concealment (selection bias)	Low risk	Sealed envelopes prepared in advance
Incomplete outcome data (attrition bias) All outcomes	High risk	Attrition partially reported; no test for differences
Selective reporting (reporting bias)	Low risk	Fully reported
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Independent assessor

Jensen 1997

Methods	RCT; 2 arms; assessed pre-treatment, post-treatment, 6 months, 18 months
Participants	End of treatment n = 59 Start of treatment n = 63
	Sex: 63 F, 0 M (at start of treatment)
	Mean age = 43.4 (SD 8.4)
	Source = pain or rehabilitation clinic
	Diagnosis = non-specific back or neck pain Mean years of pain = 4.2
	ivicali yeals of pail = 4.2
Interventions	"Woman-specific CBT"
	"Regular CBT"
Outcomes	Primary pain outcome: VAS pain intensity
	Primary disability outcome: Disability Rating Index
	Primary mood outcome: BDI depression
	Catastrophising outcome: RAI helplessness
	Pain intensity visual analogue scale
	Beck Depression Inventory (BDI) Anxiety visual analogue scale
	Disability Rating Index
	Health perception numerical rating scale
	Coping Strategies Questionnaire (CSQ)
	Rheumatoid attitudes index (helplessness)
	CBT versus active, post-treatment and follow-up: analyses 1.1, 1.2, 1.3, 2.1, 2.2, 2.3
Notes	

Jensen 1997 (Continued)

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Central randomisation using random numbers table
Allocation concealment (selection bias)	High risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	High risk	Attrition partially reported; no test for differences
Selective reporting (reporting bias)	High risk	Partially reported
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors blind to treatment condition

Jensen 2001

Methods	RCT; 4 arms; assessed at pre-treatment, post-treatment, 6 months, 18 months, 3 years
Participants	End of treatment n = 186 Start of treatment n = 214 Sex: 117 F, 93 M Mean age = 43.3 (SD 10.4) Source = pain or rehabilitation clinic Diagnosis = mixed (mostly chronic low back pain) Mean years of pain = 2.7
Interventions	"CBT" "Behavioural medicine rehabilitation" "Behaviourally orientated physical therapy" (BT) "Treatment as usual"
Outcomes	Primary pain outcome: SF36 pain (reversed) Primary disability outcome: SF36 physical function (reversed) Primary mood outcome: SF36 mental health (reversed) Catastrophising outcomes: none Short Form 36 Pain Short Form 36 Physical Function Short Form 36 Mental Health
Notes	CBT versus TAU, post-treatment and follow-up (6 months): analyses 3.1, 3.2, 3.3, 4.1, 4.2, 4.3 BT versus TAU, post-treatment and follow-up (6 months): analyses 7.1, 7.2, 7.3, 8.1, 8.2, 8.3 Baseline N used as N unavailable for post-treatment and follow-up results Yates quality scale: total quality = 27/35, design quality = 20/26, treatment quality = 7/9

Jensen 2001 (Continued)

Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Shuffled sealed envelopes
Allocation concealment (selection bias)	Low risk	Sealed envelopes; procedure by researchers blind to participant screening
Incomplete outcome data (attrition bias) All outcomes	High risk	Partially reported; differential attrition; no test of differences
Selective reporting (reporting bias)	Low risk	Fully reported
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Data gathered by research team

Kaapa 2006

Methods	RCT; 2 arms; assessed at pre-treatment, post-treatment, 6 months, 1 year, 2 years
Participants	End of treatment n = 120 Start of treatment n = 132 Sex: 120 F, 12 M (start of treatment) Mean age = 46.3 (SD 7.5) Source = community Diagnosis = chronic low back pain Mean years of pain = 1.3
Interventions	"semi-intensive multidisciplinary rehabilitation" "individual physiotherapy"
Outcomes	Primary pain outcome: pain intensity 0 to 10 Primary disability outcome: Oswestry Disability Index 0 to 100 Primary mood outcome: (DEPS) depression 0 to 30 Catastrophising outcome: none Low back pain intensity 0 to 10 Sciatic pain intensity 0 to 10 Oswestry Disability Index 0 to 100 Subjective work capacity 0 to 10 Recent sick leave due to back pain Beliefs re working (2-year follow-up) 0 to 10 The Depression Scale (DEPS) 0 to 30 Health care consumption 12 months

Kaapa 2006 (Continued)

Notes	CBT versus active, post-treatment and follow-up: analyses 1.1, 1.2, 1.3, 2.1, 2.2, 2.3
	Yates quality scale: total quality = 23/35, design quality = 20/26, treatment quality = 3/
	9

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Random numbers
Allocation concealment (selection bias)	Low risk	Opaque sealed envelopes; numbers generated by independent statistician
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Fully reported; no test for differences
Selective reporting (reporting bias)	Low risk	Fully reported
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not reported

Keefe 1990

Methods	RCT. 3 arms; assessed pre-treatment, post-treatment, 6 months
Participants	End of treatment n = 94 Start of treatment n = 99 Sex: 71 F, 28 M Mean age = 64.0 (SD 11.5) Source = rheumatology clinic Diagnosis = osteoarthritis of the knee Mean years of pain = 12.0
Interventions	"coping skills training" "arthritis education" "standard care"
Outcomes	Primary pain outcome: AIMS pain Primary disability outcome: AIMS physical disability Primary mood outcome: AIMS psychological disability Catastrophising outcome: none Arthritis Impact Measurement Scale (AIMS): pain Arthritis Impact Measurement Scale (AIMS): psychological disability Arthritis Impact Measurement Scale (AIMS): physical disability Pain behaviour (Keefe & Block) observation Coping Strategy Questionnaire

Keefe 1990 (Continued)

	Medication use
Notes	CBT versus active, post-treatment and follow-up: analyses 1.1, 1.2, 1.3, 2.1, 2.2, 2.3 CBT versus TAU, post-treatment and follow-up: analyses 3.1, 3.2, 3.3, 4.1, 4.2, 4.3 Yates quality scale: total quality = 26/35, design quality = 18/26, treatment quality = 8/9

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"randomly assigned (using a random number table)"
Allocation concealment (selection bias)	High risk	Not reported (but equal credibility of treatments rated by participants)
Incomplete outcome data (attrition bias) All outcomes	Low risk	Fully reported
Selective reporting (reporting bias)	Low risk	Fully reported
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not reported

Keefe 1996

Methods	RCT; 3 arms; assessed at pre-treatment, post-treatment, 6 months, 1 year
Participants	End of treatment n = 82 Start of treatment n = 88 Sex: 54 F, 34 M Mean age = 62.6 (SD 10.1) Source = volunteer Diagnosis = osteoarthritis of knee Mean years of pain = 10.7
Interventions	"spouse-assisted coping skills training" "coping skills training" "spouse-supported arthritis education"
Outcomes	Primary pain outcome: AIMS pain Primary disability outcome: AIMS physical disability Primary mood outcome: AIMS mental disability Catastrophising outcome: none Arthritis Impact Measurement Scale (AIMS): pain Arthritis Impact Measurement Scale (AIMS): physical

Keefe 1996 (Continued)

	Arthritis Impact Measurement Scale (AIMS): psychological Coping Strategies Questionnaire: coping Coping Strategies: pain control Pain behaviour (Keefe & Block) observation
Notes	CBT versus active, post-treatment: analyses 1.1, 1.2, 1.3 Yates quality scale: total quality = 25/35, design quality = 17/26, treatment quality = 8/9

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	"randomly assigned"
Allocation concealment (selection bias)	High risk	Not reported (but equal credibility of treatments rated by participants)
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Fully reported; no differential attrition but no test for differences
Selective reporting (reporting bias)	Low risk	Fully reported
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not reported

Kole-Snijders 1999

Methods	RCT; 3 arms; assessed at pre-treatment, post-treatment, 6 months, 1 year
Participants	End of treatment n = 133 Start of treatment n = 148 Sex: 94 F, 54 M Mean age = 30.8 (SD 9.1) Source = pain or rehabilitation clinic Diagnosis = chronic low back pain Mean years of pain = 9.8
Interventions	"operant + cognitive coping skills" "operant + group discussion" "waiting list"
Outcomes	Primary pain outcome: no data available Primary disability outcome: no data available Primary mood outcome: no data available Catastrophising outcome: none

Kole-Snijders 1999 (Continued)

	(all reduced by factor analysis to 3 scores: motoric, coping control, negative affect) Pain Behaviour Scale Checklist for Interpersonal Pain Behaviour Behavioural approach test (walking distance) Multi dimensional Locus of Control Pain Cognition Checklist Coping Strategies Questionnaire Nijmegen Hyperventilation Questionnaire Visual analogue scale: pain McGill Pain Questionnaire: pain
Notes	No data Yates quality scale: total quality = 28/35, design quality = 20/26, treatment quality = 8/9

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"independent researcher blindly drew [numbers assigned randomly to patients] and assigned to one of three conditions"
Allocation concealment (selection bias)	Low risk	independent researcher
Incomplete outcome data (attrition bias) All outcomes	Low risk	Fully reported
Selective reporting (reporting bias)	Unclear risk	Reported as factor scores
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessor unaware of treatment condition

Kraaimaat 1995

Methods	RCT; 3 arms; assessed at pre-treatment, post-treatment, 6 months
Participants	End of treatment n = 52 Start of treatment n = 58 Sex: 52 F, 25 M (from the 77 who agreed to participate) Mean age = 57.0 (SD 12.7) Source = rheumatology clinics Diagnosis = rheumatoid arthritis Mean years of pain = 15.6
Interventions	"cognitive behavioural therapy" "occupational therapy"

Kraaimaat 1995 (Continued)

	"waiting list"
Outcomes	Primary pain outcome: IRGL pain Primary disability outcome: IRGL function (Reversed) Primary mood outcome: IRGL depression Catastrophising outcome: none Invloed van Reuma op Gezondheid en Leefwijze (IRGL): function Invloed van Reuma op Gezondheid en Leefwijze (IRGL): self care Invloed van Reuma op Gezondheid en Leefwijze (IRGL): pain Invloed van Reuma op Gezondheid en Leefwijze (IRGL): anxiety Invloed van Reuma op Gezondheid en Leefwijze (IRGL): depression Invloed van Reuma op Gezondheid en Leefwijze (IRGL): potential support Invloed van Reuma op Gezondheid en Leefwijze (IRGL): actual support Invloed van Reuma op Gezondheid en Leefwijze (IRGL): mutual visits
Notes	CBT versus active, post-treatment and follow-up: analyses 1.1, 1.2, 1.3, 2.1, 2.2, 2.3 CBT versus TAU, post-treatment and follow-up: $N < 20$ Yates quality scale: total quality = $21/35$, design quality = $14/26$, treatment quality = $7/9$

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	"randomly assigned"
Allocation concealment (selection bias)	High risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Fully reported; several differences between dropouts and completers
Selective reporting (reporting bias)	Low risk	Fully reported
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not reported

Leeuw 2008

Methods	RCT; 2 arms; assessed at 2 pre-treatment occasions, post-treatment, 6-month follow-up, 12-month follow-up
Participants	End of treatment n = 77 Start of treatment n = 85 Sex: 41 F, 44 M Mean age = 45.3 (SD 9.5) Source = rehabilitation clinics, occupational health, pain department

Leeuw 2008 (Continued)

	Diagnosis = back pain (and at least moderate fear on TSK) Mean years of pain = 9
Interventions	"Exposure in vivo" "Operant graded activity"
Outcomes	Primary pain outcome: MPQ pain intensity Primary disability outcome: Quebec Back Pain Disability Scale (Dutch version) Primary mood outcome: none Catastrophising outcome: PCS 1. Quebec Back Pain Disability Scale (Dutch version) 2. Patient Specific Complaints: VAS 0 to 100 of difficulty with 3 activities 3. Perceived harmfulness of activities (PHODA) 4. Pain Catastrophizing Scale: catastrophising 5. Daily activity: actimeter 6. Pain: mean of VAS 0 to 100 scales for current, worst and least pain
Notes	December 2009 search Exposure in vivo versus operant graded activity: analyses 5.1, 5.2, 5.4, 6.1, 6.2, 6.4 Yates quality scale: total quality = 32/35, design quality = 24/26, treatment quality = 8/9

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	"predetermined and computer-generated randomization schedule"
Allocation concealment (selection bias)	Low risk	Sealed envelope; research assistant only could access randomization schedule
Incomplete outcome data (attrition bias) All outcomes	Low risk	Fully reported
Selective reporting (reporting bias)	Low risk	Fully reported
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Electronic administration of assessments

Lindell 2008

Methods	RCT; 2 arms; assessed at pre-treatment, post-treatment, 18-month follow-up
Participants	End of treatment n = 123 Start of treatment n = 125 Sex: 68 F, 57 M

Lindell 2008 (Continued)

	Mean age = 42.6 (SD not given) Source = primary care Diagnosis = non-specific back or neck pain Mean years of pain = not given but had to be sick listed for more than 6 weeks up to 2 years; mean over 7 months sick listed
Interventions	"Cognitive-behavioural rehabilitation" "Primary care"
Outcomes	Primary pain outcome: none Primary disability outcome: none Primary mood outcome: none Catastrophising outcome: none 1. Sick listed days 2. Healthcare visits
Notes	December 2009 search No data available Yates quality scale: total quality = 18/35, design quality = 16/26, treatment quality = 2/6

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computerised block randomisation procedure
Allocation concealment (selection bias)	Low risk	Randomisation generated by independent statistician; in opaque envelopes
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Fully reported; no test for differences
Selective reporting (reporting bias)	Low risk	Fully reported
Blinding of outcome assessment (detection bias) All outcomes	High risk	Assessors not blind to treatment condition, except for sick listing outcome

Litt 2009

Methods	RCT; 2 arms; CBT + standard treatment; standard treatment; post-treatment
Participants	End of treatment: n = 54 Start of treatment: n = 54 Sex: 46 F; 8 M Mean age: 41.0 (SD 11.0) Source = dental clinics and dentists (15%); newspaper and web adverts (85%)

Litt 2009 (Continued)

	Diagnosis = temporomandibular disorder Mean years of pain: 5.6 (SD 5.4)
Interventions	CBT + standard treatment; standard treatment (splint, diet, NSAIDs)
Outcomes	Primary pain outcome: MPI 0 to 6 Primary disability outcome: interference MPI 0 to 6 Primary mood outcome: CES-D Catastrophising outcome: data not available Pain Intensity MPI 0 to 6 CES-D Depression Interference with activity MPI 0 to 6 2 items modified from Catastrophising Sub-Scale CSQ Several times daily sampling of pain, control, affect, coping, catastrophising
Notes	CBT versus TAU: analyses 3.1, 3.2, 3.3 2011 update search Yates quality scale: total quality 14/35, design quality 11/26, treatment quality 3/9

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computerised urn randomisation
Allocation concealment (selection bias)	High risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	High risk	Attrition not reported
Selective reporting (reporting bias)	Low risk	Fully reported
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not reported

McCarberg 1999

Methods	RCT; 2 arms; assessed pre-treatment, 6 months follow-up
Participants	End of treatment n = 245 Start of treatment n = 353 Sex: 264 F, 89 M Mean age = 52.1 (SD 9.6) Source = pain or rehabilitation clinic Diagnosis = mixed chronic pain, many chronic low back pain Mean years of pain = 9.6

McCarberg 1999 (Continued)

Interventions	"Cognitive behaviour therapy" "minimal home study"
Outcomes	Primary pain outcome: MPI pain severity Primary disability outcome: MPI pain interference Primary mood outcome: MPI affective distress Catastrophising outcome: none 11-point box scale: pain severity Pain discomfort scale: pain distress Multidimensional Pain Inventory: pain severity Multidimensional Pain Inventory: affective distress Multidimensional Pain Inventory: self control
Notes	Multidimensional Pain Inventory: interference Multidimensional Pain Inventory: social support and spouse behaviour subscales CBT versus active, follow-up: analyses 2.1, 2.2, 2.3 Yates quality scale: total quality = 11/35, design quality = 9/26, treatment quality = 2/9

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"Patients were randomized using a computer-generated random number list"
Allocation concealment (selection bias)	High risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	No attrition during treatment, only at follow-up; no test for differences
Selective reporting (reporting bias)	Low risk	Fully reported
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not reported

Mishra 2000

Methods	RCT; 4 arms; assessed at pre-treatment, post-treatment
Participants	End of treatment n = 94 Start of treatment n = 94 Sex: 77 F, 7 M Mean age = 35.8 (SD 9.9) Source = pain or rehabilitation clinic and volunteer Diagnosis = temporomandibular joint disorder Mean years of pain = 7.0

Mishra 2000 (Continued)

Interventions	"Biofeedback" (BT) "Cognitive behavioural skills training" (CBT) "Cognitive behavioural skills training + biofeedback" "no treatment control"
Outcomes	Primary pain outcome: CPI pain index Primary disability outcome: none available Primary mood outcome: none available Catastrophising outcomes: none Characteristic Pain Index (CPI) pain severity 0 to 100 Graded Chronic Pain Score Profile of Mood States total
Notes	CBT versus TAU, post-treatment: analysis 3.1 BT versus TAU, post-treatment: analysis 7.1 Yates quality scale: total quality = 19/35, design quality = 12/26, treatment quality = 7/9

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	"patients were assigned to group in a semi- random fashion using the urn method of random assignment"
Allocation concealment (selection bias)	High risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	High risk	Attrition not reported
Selective reporting (reporting bias)	High risk	Partially reported
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not reported

Nicassio 1997

Methods	RCT; 2 arms; assessed at pre-treatment, post-treatment, 6 months
Participants	End of treatment n = 71 Start of treatment n = 96 Sex: 63 F, 8 M (at follow-up) Mean age = 53.1 (SD no given) Source = pain or rehabilitation clinic, support groups Diagnosis = fibromyalgia Mean years of pain = 11.1

Nicassio 1997 (Continued)

Interventions	"behavioural treatment" "education"
Outcomes	Primary pain outcome: not available Primary disability outcome: quality of well being Primary mood outcome: CES-D Depression Catastrophising outcome: RAI helplessness Pain index: composite of Fibromyalgia Impact Questionnaire pain scale, MPQ PRI, number of body areas, and flare index Pain Behavior Checklist self reported pain behaviour Pain behaviour (Keefe & Block) observation Center for Epidemiologic Studies Depression Scale (CES-D) Rheumatology Attitudes Index helplessness subscale Pain Management Inventory active and passive coping Quality of Well being Scale QWB: structured interview on functional impairment Quality of Social Support Scale Myalgia score, nurse rated on examination
Notes	BT versus active, post-treatment and follow-up: analyses 5.2, 5.3, 6.2, 6.3 Yates quality scale: total quality = 21/35, design quality = 15/26, treatment quality = 6/9

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	In blocks, "randomly assigned, using a random numbers table"
Allocation concealment (selection bias)	High risk	Not reported, though credibility ratings equal across treatments
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition fully reported; differential attri- tion across groups; no differences
Selective reporting (reporting bias)	Low risk	Fully reported
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not reported

Parker 1988

Methods	RCT; 3 arms; assessed at pre-treatment, 6 months, 1 year
Participants	End of treatment n = 83 Start of treatment n = not given Sex: 3 F, 80 M Mean age = 60.6 (SD 7.7) Source = hospital Diagnosis = rheumatoid arthritis Mean years of pain = 11.4
Interventions	"cognitive behavioural pain management group" "attention placebo group" "control group" (TAU)
Outcomes	Primary pain outcome: no data available Primary disability outcome: no data available Primary mood outcome: no data available Catastrophising outcome: none Visual analogue scale pain McGill Pain Questionnaire pain dimensions Coping Strategies Questionnaire Arthritis Impact Measurement Scale (AIMS) Beck Depression Inventory Symptom Checklist-90R psychological symptoms Hassles Scale Ways of Coping Questionnaire Arthritis Helplessness Index Disease status measures including walking speed
Notes	No data Yates quality scale: total quality = 17/35, design quality = 13/26, treatment quality = 4/9

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	"using a table of random numbers, subjects were assigned"
Allocation concealment (selection bias)	High risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	High risk	Attrition not reported
Selective reporting (reporting bias)	High risk	Partially reported

Parker 1988 (Continued)

В	elinding of outcome assessment (detection	High risk	Not reported
bi	ias)		
A	all outcomes		

Puder 1988

Methods	RCT; 2 arms; assessed at pre-treatment, post-treatment, 1 month
Participants	End of treatment n = 69 Start of treatment n = 71 Sex: 49 F, 20 M Mean age = 52.7 (SD 14.4) Source = community Diagnosis = mixed chronic pain Mean years of pain = 10.0
Interventions	"Cognitive behaviour therapy" "waiting list"
Outcomes	Primary pain outcome: pain diary Primary disability outcome: pain interference Primary mood outcome: none available Catastrophising outcome: none Pain diary 0 to 5: highest and lowest ratings Pain interference 0 to 5 Coping 0 to 5 Medication use
Notes	CBT versus TAU, post-treatment: analyses 3.1, 3.2 Yates quality scale: total quality = 13/35, design quality = 10/26, treatment quality = 3/9

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	"randomly assigned"
Allocation concealment (selection bias)	High risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Attrition reported; no test for differences
Selective reporting (reporting bias)	High risk	Partially reported

Puder 1988 (Continued)

Blinding of outcome assessment (detection bias) All outcomes	High risk	Not reported
Schmidt 2011		
Methods	RCT; 3 arms; mindfulness-based stress reduction, active relaxation control, waiting list; post-treatment, 2-month follow-up	
Participants	End of treatment n = 148 Start of treatment n = 177 Sex: 177 F; 0 M Mean age = 52.5 (SD 9.6) Source = newspapers, GP and specialist referrals, patient self help groups Diagnosis = fibromyalgia Mean years of pain: 4.0 (SD 3.9)	
Interventions	Mindfulness-based stress reduction; active control (relaxation, support and education); waiting list	
Outcomes	Primary pain outcome: Pain Perception Scale (sensory) Primary disability outcome: Fibromyalgia Impact Questionnaire Primary mood outcome: CES-D Catatrophising outcome: none Pain Perception Scale (Sensory and Affective) Fibromyalgia Impact Questionnaire Depression: CES-D Anxiety: Trait Sub-scale STAI Pittsburgh Sleep Quality Index Health-related Quality of Life Freiburg Mindfulness Inventory Physical symptoms: Giessen Complaint Questionnaire Ongoing therapies, medical visits and medication Medication diary Goal-attainment scaling by interview	
Notes	Active relaxation control versus waiting list; 2011 update search Yates quality scale: total quality = 31/35, de 9	·

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	"randomized in blocks by a computer algorithm"

Schmidt 2011 (Continued)

Allocation concealment (selection bias)	Low risk	Patients and personnel blinded to treatment allocation
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Attrition fully reported; no test for differences
Selective reporting (reporting bias)	Low risk	Fully reported
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors blinded

Smeets 2006

Methods	RCT; 4 arms; assessed at pre-treatment, post-treatment, 1 year
Participants	End of treatment n = 212 Start of treatment n = 223 Sex: 106 F, 117 M Mean age = 41.6 (SD 10.0) Source = pain or rehabilitation clinic Diagnosis = CLBP Mean years of pain = 4/6
Interventions	"Cognitive behavioural therapy + active physical treatment" "Cognitive behavioural therapy" "active physical treatment" "waiting list"
Outcomes	Primary pain outcome: MPQ PRI (follow-up only) Primary disability outcome: Roland & Morris Disability Scale Primary mood outcome: BDI Catastrophising outcome: process only Roland Morris Disability Questionnaire disability Difficulty with 3 most limited activities: 0 to 100 Visual analogue scale pain Beck Depression Inventory Pain Cognitions List: catastrophising, pain control subscales as process measures Follow-up only MPQ PRI 6. 5-minute walk 7. 50-foot walk 8. timed stand-to-sits 9. extended reach 10. stair climb 11. lifting task

Smeets 2006 (Continued)

Notes	1-year follow-up Smeets 2008; December 2009 search
	CBT plus active PT versus active PT: analyses 1.1, 1.2, 1.3, 2.1, 2,2. 2.3
	GA plus problem solving versus WLC: analyses 3.1, 3.2, 3.3 (waiting list not followed
	up)
	Yates quality scale: total quality = 28/35, design quality = 23/26, treatment quality = 5/
	9

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomised in blocks by computer-generated algorithm
Allocation concealment (selection bias)	Low risk	Generated by independent statistician; sealed envelopes
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Attrition fully reported; no test for differences
Selective reporting (reporting bias)	Low risk	Fully reported
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessment by blinded research assistants

Strauss 1986

Methods	RCT; 3 arms; assessed at pre-treatment, post-treatment, 6 months
Participants	End of treatment n = 43 Start of treatment n = 57 Sex: 46 F, 11 M Mean age = 54.0 (SD 13.0) Source = rheumatology clinic Diagnosis = rheumatoid arthritis Mean years of pain not given
Interventions	"group psychotherapy" "relaxation/assertion" "no treatment"
Outcomes	Primary pain outcome: no data available Primary disability outcome: no data available Primary mood outcome: no data available Catastrophising outcome: none 4 aggregate outcome measures: Functional status, social adaptation, psychological adaptation, psychological symptoms

Strauss 1986 (Continued)

	Measures contributing to these: Arthritis Impact Measurement Scale (AIMS) Short Form 36 Rathus Assertive Behavior Scale Rosenberg Self-Esteem Scale Hostility Inventory Wright's Human Service Scale & Handicap Problems Inventory
Notes	No data Yates quality scale: total quality = 10/35, design quality = 9/26, treatment quality = 1/9

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	"randomly assigned"
Allocation concealment (selection bias)	High risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	High risk	Attrition not reported
Selective reporting (reporting bias)	High risk	Partially reported
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not reported

Thieme 2003

Methods	RCT; 2 arms; assessed at pre-treatment, post-treatment, 6 months, 15 months
Participants	End of treatment n = 61 Start of treatment n = 83 Sex: 61 F, 0 M Mean age = 47.3 (SD 8.3) Source = hospital for rheumatic disorders Diagnosis = fibromyalgia Mean years of pain = 16.5
Interventions	"operant treatment" "standard physical treatment"
Outcomes	Primary pain outcome: MPI pain Primary disability outcome: MPI interference Primary mood outcome: MPI affective distress Catastrophising outcome: none

Thieme 2003 (Continued)

	Diary pain intensity	
	, 1	
	Multidimensional Pain Inventory: pain	
	Multidimensional Pain Inventory: interference	
	Multidimensional Pain Inventory: life control	
	Multidimensional Pain Inventory: affective distress	
	Multidimensional Pain Inventory: social support	
	Multidimensional Pain Inventory: self efficacy	
	Multidimensional Pain Inventory: punishing responses, solicitous responses, distracting	
	responses	
	Multidimensional Pain Inventory: total activities	
	Doctor visits (from medical records)	
	Hospital days (from medical records)	
	Sleep hours diary	
	Medication diary	
	Tubingen pain behaviour scale	
Notes	BT versus TAU, post-treatment and follow-up: analyses 7.1, 7.2, 7.3, 8.1, 8.2, 8.3 Yates quality scale: total quality = 15/35, design quality = 11/26, treatment quality = 4/9	

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	"randomly assigned"
Allocation concealment (selection bias)	High risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Attrition reported; no test for differences
Selective reporting (reporting bias)	Low risk	Fully reported
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not reported

Thorsell 2011

Methods	RCT; 2 arms; self help acceptance and commitment therapy, self help applied relaxation; post-treatment: 6-month and 12-month follow-up
Participants	End of treatment: n = 64 Start of treatment: n = 98 Sex: 63 F; 35 M Source = pain clinic Diagnosis = mixed chronic pain

Thorsell 2011 (Continued)

	Mean age: 46.0 (SD 12.3) Mean years of pain: not given (98% more than 1 year)
Interventions	Self help acceptance and commitment therapy; self help applied relaxation
Outcomes	Primary pain outcome: pain intensity 0 to 10 Primary disability outcome: OMPQ 5 items Primary mood outcome: Depression HADS Catastrophising outcome: none Pain intensity 0 to 10 Function: 5 items 0 to 10 from Orebro Musculoskeletal Pain Questionnaire (reverse direction) Depression HADS Anxiety HADS Satisfaction With Life Scale Chronic Pain Acceptance Questionnaire
Notes	ACT versus active control: analyses 1.1, 1.2, 1.3, 2.1, 2.2, 2.3 2011 update search Yates quality scale: total quality 18/35, design quality 13/26, treatment quality 5/9

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"randomized by drawing pieces of paper with type of intervention"
Allocation concealment (selection bias)	High risk	Not reported, but treatment credibility equal
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Attrition fully reported; no test for differences
Selective reporting (reporting bias)	Low risk	Fully reported
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not reported

Turner 1988

Methods	RCT; 3 arms; assessed at pre-treatment, post-treatment, 6 months, 1 year
Participants	End of treatment n = 53 Start of treatment n = 81 Sex: 30 F, 51 M Mean age = 46.0 (SD not given)

Turner 1988 (Continued)

	Source = pain or rehabilitation clinic Diagnosis = CLBP Mean years of pain = 6.2
Interventions	"CBT" "operant behavior therapy" "waiting list"
Outcomes	Primary pain outcome: MPQ PRI Primary disability outcome: SIP patient-rated Primary mood outcome: none available Catastrophising outcome: CEQ Multidimensional Pain Questionnaire: Pain Response Index Sickness Impact Profile: patient-rated Sickness Impact Profile: spouse-rated Pain behaviour (Keefe & Block) observation Pain Behavior Checklist patient-rated Pain Behavior Checklist spouse-rated Cognitive Errors Questionnaire
Notes	CBT versus TAU, post-treatment (waiting list not followed up): analyses 3.1, 3.2 BT versus TAU, post-treatment (waiting list not followed up): analyses 7.2 Yates quality scale: total quality = 23/35, design quality = 15/26, treatment quality = 8/9

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	"randomly assigned"
Allocation concealment (selection bias)	High risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Attrition fully reported; no test for differences
Selective reporting (reporting bias)	Low risk	Partially reported but full account of excluded measures
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not reported

Turner 2006

Methods	RCT.; 2 arms; assessed at pre-treatment, post-treatment, 6 months, 1 year
Participants	End of treatment n = 142 Start of treatment n = 158 Sex: 128 F, 30 M Mean age = 37.4 (SD 11.3) Source = pain or rehabilitation clinic Diagnosis = temporomandibular joint pain Mean years of pain = not given
Interventions	"brief CBT: Pain Management Training" "education/attention control: Self care control"
Outcomes	Primary pain outcome: Graded Chronic Pain Scale: Pain Intensity Primary disability outcome: none available Primary mood outcome: BDI depression Catastrophising outcome: PCS Graded Chronic Pain Scale: Activity Interference Graded Chronic Pain Scale: Pain Intensity Beck Depression Inventory (BDI) Mandibular Function Impairment Questionnaire (MFIQ) Survey of Pain Attitudes (SOPA) TMD self efficacy scale CSQ catastrophising subscale Pain Catastrophizing Scale rumination subscale Chronic Pain Coping Inventory (CPCI) task persistence, coping self statements, relaxation, rest
Notes	CBT versus active, post-treatment and follow-up: analyses 1.1, 1.3, 2.1, 2.3 Yates quality scale: total quality = 27/35, design quality = 22/26, treatment quality = 5/9

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated by biostatistician
Allocation concealment (selection bias)	Low risk	Sealed envelopes; independent personnel; treatment credibility unequal so used as covariate
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Attrition fully reported; no test for differences
Selective reporting (reporting bias)	Low risk	Fully reported

Turner 2006 (Continued)

Blinding of outcome assessment (detection bias)	High risk	Not reported
All outcomes		

Van Koulil 2010

Methods	RCT; 2 arms; CBT: WLC; post-treatment: 6-month follow-up
Participants	End of treatment: n = 152 Start of treatment: n = 158 Sex: 148 F, 10 M Mean age: 40.8 (SD 10.5) Mean years of pain: not given (< 5 years since diagnosis) Source = rheumatology clinics Diagnosis = fibromyalgia
Interventions	Tailored CBT with exercise training; waiting list control
Outcomes	Primary pain outcome: Pain IRGL Primary disability outcome: Mobility IRGL Primary mood outcome: Negative mood IRGL Catastrophising outcome: none Pain: 6 items of IRGL Disability: 7 mobility items of IRGL (reversed) Impact: Fibromyalgia Impact Questionnaire Negative mood: 6 items of IRGL Anxiety: 10 items of IRGL
Notes	CBT versus WLC: analyses 3.1, 3.2, 3.3, 4.1, 4.2, 4.3 2011 update search Yates quality scale: total quality 24/35, design quality 15/26, treatment quality 9/9

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	"randomized in clusters"
Allocation concealment (selection bias)	High risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Attrition reported; 2 differences between dropouts and completers
Selective reporting (reporting bias)	Low risk	Fully reported

Van Koulil 2010 (Continued)

Blinding of outcome assessment (detection bias)	High risk	Not reported
All outcomes		

Vlaeyen 1996

Methods	RCT; 3 arms; assessed at pre-treatment, post-treatment, 6 months, 1 year
Participants	End of treatment $n = 122$
	Start of treatment $n = 131$
	Sex: 110 F, 15 M
	Mean age = 44.0 (SD 9.4)
	Source = pain or rehabilitation clinic
	Diagnosis = fibromyalgia
	Mean years of pain = 10.2
Interventions	"cognitive + group discussion"
	"education + group discussion"
	"waiting list"
Outcomes	Primary pain outcome: pain intensity score
	Primary disability outcome: none available
	Primary mood outcome: BDI depression
	Catastrophising outcome: none
	Composite scores from factor analysis:
	Pain intensity, pain coping, pain control, relaxation, catastrophising, pain behaviour,
	activity
	Measures contributing to factors:
	Multidimensional Pain Questionnaire: Pain Response Index
	Coping Strategies Questionnaire (CSQ)
	Beck Depression Inventory (BDI) (none available)
	Fear Survey Schedule
	Arthritis knowledge
	Maudsley Obsessive Compulsive Inventory
	Pain behaviour scale
	Multidimensional Pain Locus of Control Scale (MPCL)
	Walking distance, walking time, cycling time
Notes	CBT versus active, post-treatment: analyses 1.1, 1.3
	Yates quality scale: total quality = 20/35, design quality = 16/26, treatment quality = 4/
	9

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	"randomly assigned"

Vlaeyen 1996 (Continued)

Allocation concealment (selection bias)	High risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition reported
Selective reporting (reporting bias)	Low risk	Fully reported
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not reported

Wetherell 2011

Bias

Methods	RCT; 2 arms; acceptance and commitment therapy, CBT; post-treatment and 6 month follow-up
Participants	End of treatment: n = 99 Start of treatment: n = 114 Sex: 58 F; 56 M Mean age: 54.9 (SD 12.5) Mean years of pain: 15 (SD 35.5) Source = primary care (40%); adverts and newspaper article (40%); pain support groups (10%); various (10%) Diagnosis = mixed chronic pain
Interventions	ACT versus CBT
Outcomes	Primary pain outcome: BPI pain severity Primary disability outcome: BPI interference Primary mood outcome: BDI Catastrophising outcome: none Pain severity: BPI Sub-scale Disability: BPI Interference Sub-scale (primary outcome) Disability: MPI General Activity Sub-scale Depression: BDI-II Anxiety: PASS Quality of life: SF-12 physical and mental subscores Treatment Satisfaction Questionnaire
Notes	ACT versus CBT: analyses 1.1, 1.2, 1.3, 2.1, 2.2, 2.3 2011 update search. Yates quality scale: total quality = 32/35, design quality = 24/26, treatment quality = 8/9
Risk of bias	

Authors' judgement

Support for judgement

Wetherell 2011 (Continued)

Random sequence generation (selection bias)	Low risk	Group randomisation generated by computer
Allocation concealment (selection bias)	Low risk	Staff member who accessed randomisation code had no contact with participants
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Attrition fully reported; several differences between dropouts and completers
Selective reporting (reporting bias)	Low risk	Fully reported
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessment staff blind to treatment condition

Williams 1996

Methods	RCT; 3 arms; assessed at pre-treatment, post-treatment, 6 months, 1 year
Participants	End of treatment n = 99 Start of treatment n = 121 Sex: 68 F, 53 M Mean age = 50.0 (SD 11.5) Source = pain clinic Diagnosis = mixed chronic pain, low back commonest Mean years of pain = 7.8
Interventions	"inpatient CBT" "outpatient CBT" "waiting list"
Outcomes	Primary pain outcome: VAS pain Primary disability outcome: SIP patient-rated Primary mood outcome: BDI depression Catastrophising outcome: CSQ catastrophising Visual analogue scale (VAS): pain intensity Visual analogue scale (VAS): pain distress Sickness Impact Profile (SIP): patient-rated Beck Depression Inventory (BDI) State-Trait Anxiety Inventory (STAI) Coping Strategies Questionnaire (CSQ): catastrophising Pain Self-Efficacy Questionnaire (PSEQ) Pain Cognitions Questionnaire (PCQ) Walk distance Arm endurance Stair climb Stand ups

Williams 1996 (Continued)

	Medication use Health care use
Notes	CBT versus TAU, post-treatment (waiting list not followed up): analyses 3.1, 3.2, 3.3 Yates quality scale: total quality = 22/35, design quality = 15/26, treatment quality = 7/9

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"Randomly assigned by throw of a die"
Allocation concealment (selection bias)	High risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition reported
Selective reporting (reporting bias)	High risk	Partially reported
Blinding of outcome assessment (detection bias) All outcomes	Low risk	"interviewers and assistants blind to the patients' treatment"

Zautra 2008

Methods	RCT; 3 arms; Assessed at pre-treatment, post-treatment, 6 months follow-up
Participants	Start of treatment N = 142 End of treatment N = 137 46 M, 97 F Mean age 62.1 men, 50.6 women Diagnosis = rheumatoid arthritis Mean years of rheumatoid arthritis 15.4 years men, 11.6 years women
Interventions	"cognitive behavioral therapy for pain" "mindfulness meditation and emotion regulation therapy" "education-only group"
Outcomes	Primary pain outcome: pain diary 0 to 100 Primary disability outcome: none Primary mood outcome: PANAS negative affect Catastrophising outcome: CSQ catastrophising subscale rescored Pain once-daily diary 0 to 100 Positive and Negative Affect Schedule (PANAS): provides positive affect and negative affect scores Depressive symptoms: sum of 6 items

Zautra 2008 (Continued)

	Pain coping efficacy (2 items, 1 to 5) CSQ catastrophising subscale Pain control 1 to 10 Disease Activity Score from examination of 28 joints by rheumatologist Interleukin IL-6
Notes	December 2009 search Data obtained from author Used CBT for pain and education control group: 1.1, 1.3, 1.4, 2.1, 2.3, 2.4 Yates quality scale: total quality = 27/35, design quality = 19/26, treatment quality = 8/9

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Random numbers table
Allocation concealment (selection bias)	High risk	Not reported; treatment credibility measured but at end of treatment
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Attrition fully reported; no test for differences
Selective reporting (reporting bias)	Low risk	Fully reported
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessment by staff not involved in treatment

AIMS: Arthritis Impact Measurement Scale; BDI: Beck Depression Inventory; BT: behaviour therapy; CBT: cognitive behavioural therapy; CEQ: Cognitive Errors Questionnaire; CES-D: Center for Epidemiologic Studies Depression Scale; CLBP: chronic low back pain; CSQ: Coping Strategies Questionnaire; DASS: Depression, Anxiety & Stress Scale; EMG: electromyograph; FESV: Pain-Related Distress Questionnaire; FIQ: Fibromyalgia Impact Questionnaire; GA: graded activity; HADS: Hospital Anxiety and Depression Scale; HSCL: Hopkins Checklist; IRGL: Invloed van Reuma op Gezondheid en Leefwijze; MPQ PRI: Melzack Pain Questionnaire Pain Response Index; NRS: numerical rating scale; OMPQ: Orebro Musculoskeletal Pain Questionnaire; PANAS: Positive and Negative Affect Schedule; PCCL: Pain Coping and Cognition List; PCS: Pain Catastrophizing Scale; PDI: Pain Disability Index; PRSS: Pain-Related Self-Statements; PT: physical treatment; RAI: Rheumatoid Arthritis Index; RCT: randomised controlled trial; SD: standard deviation; SIP: Sickness Impact Profile; SLE: systemic lupus erythematosus; SOPA: Survey of Pain Attitudes; TAU: treatment as usual; TSK: Tampa Scale for Kinesiophobia; VAS: visual analogue scale; WHO: World Health Organization; WHYMPI: West Haven Yale Multidimensional Pain Inventory; WLC: waiting list control.

Characteristics of excluded studies [ordered by study ID]

Study	Reason for exclusion
Abbott 2010	Insufficient psychotherapeutic content
Abrahamsen 2008	Hypnosis study
Appelbaum 1988	Inadequate n: the number of patients in any treatment arm was less than 10
Asenlof 2005	Not chronic pain
Astin 2003	Insufficient psychotherapeutic content
Babu 2007	N < 20
Becker 2000	Insufficient psychotherapeutic content (participants could opt out of psychology and 71% did)
Bendix 1997	Insufficient psychotherapeutic content
Bradley 1987	N < 20
Broderick 2004	Insufficient psychotherapeutic content
Brox 2003	Insufficient psychotherapeutic content
Buhrman 2004	Internet trial
Carson 2005	Insufficient psychotherapeutic content
Carson 2010	Insufficient psychotherapeutic content
Castel 2009	n < 10
Christiansen 2010	Not all had chronic pain
Cook 1998	N < 20
Corrado 2003	No primary psychological treatment for pain or non-psychological comparator
Currie 2000	No primary psychological treatment for pain or non-psychological comparator
Dahl 2004	N < 10
Dalton 2004	Not chronic pain
de Sousa 2009	Insufficient psychotherapeutic content
Dufour 2010	Insufficient psychotherapeutic content

Dworkin 1994 Intervention pre-dental procedure: no outcome of psychology intervention available Dworkin 2002a Insufficient psychotherapeutic content Edinger 2005 Insufficient psychotherapeutic content Edinger 2005 No primary psychological treatment for pain or non-psychological comparator Ersek 2003 N < 20 Esmer 2010 Insufficient psychotherapeutic content Evans 2003 Not chronic pain Fairbank 2005 Cross-over trial and data on outcome collected after cross-over Ferrari 2006 Not clearly randomised Flor 1993 N < 20 Fors 2000 Insufficient psychotherapeutic content Freeman 2002 Insufficient psychotherapeutic content Garcia-Campayo 2009 Trial plan not trial George 2008 Insufficient psychotherapeutic content Glombiewski 2010a Not a treatment trial Haugstad 2006 Insufficient psychotherapeutic content Jensen 2009 Hypnosis study Johansson 1998 N < 20 Kapitza 2010 Insufficient psychotherapeutic content Keefe 2004 N < 20 Keller 2004 Insufficient psychotherapeutic content Kerns 1986 N < 10 Kroenke 2009 Insufficient psychotherapeutic content Lanb 2010 Insufficient psychotherapeutic content Lanb 2010 Insufficient psychotherapeutic content		
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Kerns 1986 N < 10 Kroenke 2009 Insufficient psychotherapeutic content	Keefe 2004	N < 20
Kroenke 2009 Insufficient psychotherapeutic content	Keller 2004	Insufficient psychotherapeutic content
	Kerns 1986	N < 10
Lamb 2010 Insufficient psychotherapeutic content	Kroenke 2009	Insufficient psychotherapeutic content
	Lamb 2010	Insufficient psychotherapeutic content

Lambeek 2009	Insufficient psychotherapeutic content
Li 2006	Insufficient psychotherapeutic content
Liedl 2011	N < 20
Linton 1984	N < 10
Linton 1985	N < 10
Linton 2001	Not chronic pain
Linton 2005	Not chronic pain
Linton 2008	N < 20
Lorig 2008	Internet trial
Machado 2007	Insufficient psychotherapeutic content (counselling)
Marhold 2001	N < 20
Menzel 2006	N < 10
Moffett 2005	Insufficient psychotherapeutic content
Moore 1985	N < 20
Moore 2000	Not chronic pain
Morone 2008	Insufficient psychotherapeutic content
Morone 2009	Insufficient psychotherapeutic content
Newton-John 1995	N < 20
Nicholas 1991	N < 10
Nicholas 1992	N < 10
O'Leary 1988	N < 20
Parker 2003	Intervention for depression not pain
Peters 1990	N < 10
Radojevic 1992	N < 20

Redondo 2004	N < 20
Rendant 2011	Insufficient psychotherapeutic content
Sahin 2011	Insufficient psychotherapeutic content
Schulze 2008	Not random allocation
Schweikert 2006	Insufficient psychotherapeutic content
Sharpe 2001	Not chronic pain
Smeets 2009	Study of predictors not outcomes of intervention
Soderlund 2001	Insufficient psychotherapeutic content
Spence 1989	N < 20
Spence 1995	N < 20
Strong 1998	Insufficient psychotherapeutic content
Turner 1982	N < 10
Turner 1990	N < 20
Turner 1993	N < 20
Turner 2011	Insufficient psychotherapeutic content
Turner-Stokes 2003	Equivalence trial
Van den Hout 2003	Not chronic pain
Van Lankveld 2004	No primary psychological treatment for pain or non-psychological comparator
Vlaeyen 1995	N < 20
Wicksell 2008	N < 20
Wong 2011	Insufficient psychotherapeutic content
Woods 2008	N < 20

Characteristics of studies awaiting assessment [ordered by study ID]

Bergdahl 1995

Methods	RCT; 2 arms; CT and "attention control" equivalent to treatment as usual
Participants	End of treatment: n = 30 Start of treatment: n = 30 Sex: 24 F; 6 M Mean age: 46 (range 38 to 57) Mean years of pain: not given Source: not given Diagnosis: resistant burning mouth syndrome
Interventions	Cognitive therapy; regular monitoring
Outcomes	Pain on 1 to 7 scale
Notes	Not identified by electronic searches but from references of another review

DATA AND ANALYSES

Comparison 1. Cognitive behavioural vs active control post-treatment

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Pain	13	1258	Std. Mean Difference (IV, Random, 95% CI)	-0.10 [-0.24, 0.04]
2 Disability	12	1130	Std. Mean Difference (IV, Random, 95% CI)	-0.19 [-0.33, -0.05]
3 Mood	13	1256	Std. Mean Difference (IV, Random, 95% CI)	-0.05 [-0.19, 0.09]
4 Catastrophising	6	735	Std. Mean Difference (IV, Random, 95% CI)	-0.18 [-0.36, 0.00]

Comparison 2. Cognitive behavioural vs active control follow-up

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Pain	11	1261	Std. Mean Difference (IV, Random, 95% CI)	-0.08 [-0.23, 0.06]
2 Disability	12	1295	Std. Mean Difference (IV, Random, 95% CI)	-0.15 [-0.28, -0.02]
3 Mood	11	1261	Std. Mean Difference (IV, Random, 95% CI)	-0.07 [-0.18, 0.05]
4 Catastrophising	2	282	Std. Mean Difference (IV, Random, 95% CI)	0.06 [-0.18, 0.29]

Comparison 3. Cognitive behavioural vs treatment as usual

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Pain	16	1148	Std. Mean Difference (IV, Random, 95% CI)	-0.21 [-0.37, -0.05]
2 Disability	15	1105	Std. Mean Difference (IV, Random, 95% CI)	-0.26 [-0.47, -0.04]
3 Mood	12	899	Std. Mean Difference (IV, Random, 95% CI)	-0.38 [-0.57, -0.18]
4 Catastrophising	5	308	Std. Mean Difference (IV, Random, 95% CI)	-0.53 [-0.76, -0.31]

Comparison 4. Cognitive behavioural vs treatment as usual follow-up

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Pain	7	635	Std. Mean Difference (IV, Random, 95% CI)	-0.09 [-0.25, 0.08]
2 Disability	6	450	Std. Mean Difference (IV, Random, 95% CI)	-0.13 [-0.51, 0.25]
3 Mood	7	637	Std. Mean Difference (IV, Random, 95% CI)	-0.26 [-0.51, -0.00]
4 Catastrophising	1		Std. Mean Difference (IV, Random, 95% CI)	Totals not selected

Comparison 5. Behavioural vs active control post-treatment

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Pain	1		Std. Mean Difference (IV, Random, 95% CI)	Totals not selected
2 Disability	2	148	Std. Mean Difference (IV, Random, 95% CI)	-0.26 [-0.58, 0.07]
3 Mood	1		Std. Mean Difference (IV, Random, 95% CI)	Totals not selected
4 Catastrophising	2	146	Std. Mean Difference (IV, Random, 95% CI)	-0.28 [-0.60, 0.05]

Comparison 6. Behavioural vs active control follow-up

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Pain	1		Std. Mean Difference (IV, Random, 95% CI)	Totals not selected
2 Disability	2	144	Std. Mean Difference (IV, Random, 95% CI)	-0.17 [-0.50, 0.16]
3 Mood	1		Std. Mean Difference (IV, Random, 95% CI)	Totals not selected
4 Catastrophising	1		Std. Mean Difference (IV, Random, 95% CI)	Totals not selected

Comparison 7. Behavioural vs treatment as usual post-treatment

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Pain	5	484	Std. Mean Difference (IV, Random, 95% CI)	-0.27 [-0.79, 0.24]
2 Disability	5	504	Std. Mean Difference (IV, Random, 95% CI)	-0.41 [-0.98, 0.16]
3 Mood	3	278	Std. Mean Difference (IV, Random, 95% CI)	-0.53 [-1.42, 0.35]
4 Catastrophising	3	269	Std. Mean Difference (IV, Random, 95% CI)	-0.72 [-1.43, -0.01]

Comparison 8. Behavioural vs treatment as usual follow-up

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Pain	2	182	Std. Mean Difference (IV, Random, 95% CI)	-0.03 [-0.32, 0.26]
2 Disability	3	336	Std. Mean Difference (IV, Random, 95% CI)	-0.54 [-1.51, 0.44]
3 Mood	2	160	Std. Mean Difference (IV, Random, 95% CI)	-0.65 [-2.07, 0.77]

WHAT'S NEW

Last assessed as up-to-date: 10 September 2012.

Date	Event	Description
27 July 2017	Amended	Author deceased. See Published notes.
9 February 2016	Review declared as stable	See Published notes.

HISTORY

Protocol first published: Issue 4, 2008 Review first published: Issue 2, 2009

Date	Event	Description
23 March 2016	Amended	Amended declarations of interest section (see Declarations of interest).
19 December 2012	Amended	Minor correction to the PLS.
13 July 2012	New search has been performed	We included 12 new trials from two new searches (Bliokas 2007; Ehrenborg 2010; Glombiewski 2010b; Leeuw 2008; Lindell 2008; Litt 2009; Morone 2008; Schmidt 2011; Thorsell 2011; Van Koulil 2010; Wetherell 2011; Zautra 2008). Thirty four trials included in the previous version were excluded (Astin 2003; Babu 2007; Becker 2000; Bradley 1987; Buhrman 2004; Carson 2005; Cook 1998; Dworkin 1994; Dworkin 2002b; Ersek 2003; Fairbank 2005; Flor 1993; Freeman 2002; Johansson 1998; Keefe 2004; Linton 2008; Marhold 2001; Moore 1985; Newton-John 1995; O'Leary 1988; Peters 1990; Radojevic 1992; Redondo 2004; Spence 1989; Spence 1995; Strong 1998; Turner 1990; Turner 1993; Turner-Stokes 2003; Vlaeyen 1995; Wicksell 2008; Woods 2008). We raised the criterion for entry from N>10 to N>20 in each arm. We added 'Risk of bias' ratings for all included studies. We also added a new outcome: catastrophic thinking
29 March 2012	New citation required and conclusions have changed	The evidence for CBT is stronger, particularly when compared with treatment as usual/waiting list, and for mood and catastrophic thinking. The evidence for behaviour therapy is weak or lacking. The field will not

be further advanced by more small RCTs of variants of CBT for heterogeneous patient groups but by different trial and analytic methods

CONTRIBUTIONS OF AUTHORS

AW oversaw the review and authoring of the manuscript, and authored sections of the manuscript. AW, SM and CE all authored sections of the manuscript, extracted data from papers and made quality ratings. SM advised on statistical strategy. All authors contributed to conceptualisation of the review, selection of papers and judging the quality of the studies.

DECLARATIONS OF INTEREST

Following discussions with the Cochrane Funding Arbiter in 2015/16, we have revised and expanded our declarations of interest to fully comply with the updated Cochrane Commercial Sponsorship Policy (see http://community-archive.cochrane.org/organisational-policy-manual/appendix-5-commercial-sponsorship-policy).

ACdeCW: UCL London received payment from Astellas Pharma Europe for her to speak about the psychology of pain at a general pain meeting in 2015. She is an author of an included study but was not involved in the data extraction or ratings of bias and quality.

CE attended a meeting of IMMPACT in 2011, an organisation that develops outcome measures and consults on analgesic trial design. IMMPACT receives arm's length funding from numerous pharmacological, charitable, and Governmental bodies (including the FDA). Research funding unrelated to this study was received by the University of Bath Centre for Pain Research from Reckitt Benckiser Healthcare during this review production.

SJM: none known.

SOURCES OF SUPPORT

Internal sources

• No sources of support supplied

External sources

• Department of Health, UK. Incentive Scheme Grant

DIFFERENCES BETWEEN PROTOCOL AND REVIEW

- 1. No data were available in the trials on adverse events, withdrawal and escape or emergency analgesia.
- 2. No dichotomous outcomes were reported so no numbers needed to treat (NNTs) were calculated.
- 3. No adjustment for reliability of measures was made.
- 4. Planned subgroup analyses on doses and on different conditions were not undertaken due to lack of data.
- 5. The criterion in the protocol of a minimum of 10 participants in each arm for entry into analyses was raised to a minimum of 20, given the demonstrated association between small numbers and bias (Ioannidis 2005; Moore 2010; Nuesch 2009).
- 6. A new outcome variable, catastrophic thinking, was included for all contrasts. This has emerged as a predictor of behavioural and emotional outcomes in the longer term, and is a widely (if not universally) used target of cognitive treatment.
- 7. Assessment of risk of bias in included studies: this has been expanded to include a fuller description, using the *Cochrane Handbook* recommendations.
- 8. Data extraction for 46 of the 60 trials in our penultimate selection (77%) was done independently by two authors, and the remainder by one.

NOTES

This is an active area of development but at February 2016 there were no new potentially relevant studies likely to change the conclusions. Therefore, this review has now been stabilised following discussion with the authors and editors. The review will be re-assessed for updating in 2021.

Author Stephen Morley sadly passed away in 2017. The review has been republished in July to reflect this.

INDEX TERMS

Medical Subject Headings (MeSH)

Affect; Behavior Therapy [*methods]; Chronic Pain [psychology; *therapy]; Cognitive Therapy [methods]; Randomized Controlled Trials as Topic

MeSH check words

Adult; Humans